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(54) **Balloon catheter for occluding aneurysms or branch vessels**

Ballonkatheter zum Verschliessen von Aneurysmen oder Nebenblutgefässen

Cathéter à ballon d'obturation pour anéurismes ou vaisseaux latéraux

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(56) References cited:
DE-A- 3 227 575 **US-A- 4 351 342**
US-A- 4 832 688 **US-A- 5 041 090**

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Description

The present invention relates to apparatus for delivering occluding agents through the opening in a blood vessel wall and into the aneurysm chamber or into a peripheral vessel, and in particular by introducing a balloon catheter, inflating the balloon to seal the blood vessel lumen around the vessel opening, and delivering the occluding agent through the opening to prevent loss of occluding agent into the blood vessel during such delivery and until in situ stabilization of the occluding agent as an occluding cast shaped to retain the patency of the blood vessel at the occluded opening.

The occurrence of aneurysms in weakened blood vessel walls, particularly in arterial blood vessels, often presents a life threatening risk to a patient. This is particularly true in blood vessels serving the heart, brain and other vital organs. In both venous and arterial blood vessels, such aneurysms may rupture, causing internal bleeding and loss of blood pressure or become the source of clots that may become dislodged and are borne by moving blood to other sites where they restrict blood flow. In major arteries, the rupture may lead to severe loss of pressure and rapid death. In the brain, the pooling of blood may lead to pressure on brain cells and result in a stroke.

The invasive, surgical removal of aneurysms and closure of the opening in the vessel wall itself presents a grave risk to the patient and poses severe post-operative complications. Conventional vascular surgery or micro-surgery may be employed successfully to correct aneurysms of vessels accessible to such surgical approach. However, the most threatening aneurysms are often deep within a vital organ or large in size and having a wide neck where the trauma of surgical treatment presents a great risk.

One surgical treatment for dealing with such aneurysms involves closing off the blood vessel and thereby sacrificing it and tissue that it serves to ischemia. This type of occlusion may be employed as a last resort in cases where the induced trauma presents the lesser evil.

It is therefore often either impossible to proceed surgically or preferable to avoid such surgical procedures. Less invasive approaches have been proposed and tested either in animals or clinically for closing off the aneurysm opening and/or filling the aneurysm sac. For example, as described in U. S. Patent No. 5,041,090, a number of approaches have been undertaken employing catheters for positioning detachable balloons either in the aneurysm sac or in the adjacent vessel, filling the balloon with a quick setting polymer and detaching the balloon in attempts to either fill and displace blood in the aneurysm sac or effect stationary occlusion of the vessel. In the '090 patent, a pair of catheter borne, detachable balloons are provided that are intended to be placed in the aneurism sac, inflatable with one balloon inside the other, and inflated to fill with

one material or polymerize another material in the outer balloon in a shape conforming to the aneurism sac, so that the catheters may be detached with the balloons remaining in place. In one embodiment, polymerizable material appears to be directed out of multiple holes in the outer balloon and into contact with the wall of the aneurism to bond the balloons and wall together.

In all of these procedures, it is necessary to get a balloon or balloons directed through the neck of the aneurism and retained there. The inflated balloon or balloons may protrude out through the opening of the aneurism and interfere with blood flow through the vessel lumen. Material emitted from holes in a balloon in the aneurism chamber may be forced out the opening and block the vessel lumen. Moreover, the introduction of the balloon catheter or catheters into the aneurism chamber itself poses the risk that the aneurism wall will be punctured.

Further approaches to encouraging thrombus formation in aneurism chambers are described in U.S. Patent No. 5,234,437 wherein a plurality of metallic vaso-occlusion coils are placed in the chamber by positioning a pusher catheter into the opening and detaching the coils. Detachment is proposed by applying current of electrically sever the connection or by threading the coil out of engagement with the pusher mechanism. See also, for example, "Electro Thrombosis of Aneurysms" *J. Neurosurgery*, 75:1-7, 1/91 by Guglielmo, where electric current is employed as well. These approaches require the proper introduction of a catheter into the opening, and the positioning of such catheters is difficult and presents the risk of perforating the aneurism wall.

In a still further approach described in U.S. Patent No. 5,219,355, a catheter borne sleeve is proposed to be placed across and within the depicted wide mouth opening of an aneurism to block the mouth and serve as an alternative, intraluminal blood vessel. The sleeve is attached at either end to a pair of expandable stents which are introduced by a double balloon catheter and expanded upon inflation of the balloons. Presumably, the expanded stents stabilize the sleeve ends against patent blood vessel walls. In this approach, an apparent risk lies in detachment of one or both of the stents from contact with the blood vessel walls over time. Alternatively, as the stents fibrose in, the fibrosis may restrict blood flow and lead to further complications. Many blood vessels would appear to be too small to benefit from this approach due to the necessary size of the components and introducing apparatus.

Spaced apart, double balloon catheters are also proposed for use in temporarily occluding blood vessels to introduce a therapeutic agent in treating blood vessel intima injured in balloon angioplasty procedures, as disclosed in U.S. Patent No. 4,832,688.

Further spaced apart balloon catheters have been proposed for isolating a network of small branch blood vessels collaterally supplying a tumor and injecting a

contrast material or a small vessel occlusive collagen material into the network, as disclosed in U.S. Patent Nos. 4,655,746 and 4,708,718. The injected material is delivered directly into the vessel between the two inflated balloons and described as being drawn into the small diameter branch blood vessels and remaining there over a period of time to occlude them. The disposal of the collagen material remaining in the main vessel is apparently not explained. To the extent that the solidified collagen remains in the secondary and tertiary vessels after withdrawal of the catheter, it may protrude out of these minor branch vessels into the lumen of the main vessel and provide sites for the development of thrombi or stenosis.

DE-A-3227575 which is used as basis for the preamble of claim 1 discloses another spaced apart balloon catheter for delivery an occluding agent.

In addition, single or co-axially disposed double balloon catheters of the type disclosed in U.S. Patent Nos. 5,049,132, 5,087,244, 5,112,305, and 5,213,576 are described for distributing therapeutic agents through side wall holes in the outermost balloon to vessel walls to treat an atherosclerotic plaque or to induce penetration of the agent into the vessel wall to treat a vessel wall tumor or for applying heparin post-operatively at the site of an angioplasty procedure. These balloon catheters infuse therapeutic agents into the vessel itself and are not suited to the introduction of an occluding agent of a type that would also occlude an aneurysm chamber or branch vessel.

In a further surgical procedure for harvesting saphenous veins for use in bypass surgery, it is known to strip out the saphenous vein section, remove the vein valves and tie off the branching vessels to prepare the section for implant as a bypass artery section. It is also known to do an in situ bypass converting a section of a vein adjacent a blocked artery section into a substitute artery section. The vein section is tied off proximally and distally, severed, and the two ends are grafted to the artery. In order to prepare the vein section, it is necessary to excise any venous valves in the vein section lumen and to tie off any peripheral or side branch veins. See, for example, "Endo-Vascular Infrainguinal In Situ Saphenous Bypass: A Multi Center Report", *J. Vascular Surgery*, 1992;16:453-458, by Rosenthal.

To avoid the surgery to expose the length of the sacrificed vein section to tie off the side branches, it has been proposed steer a catheter into the side branches and deposit occlusion coils therein to occlude the vessels.

Despite the advances and improvements in treatments for various conditions that have been introduced in recent years through the use of balloon catheters, a need remains for an apparatus for intraluminally occluding aneurysms in a main blood vessel wall that is simple to practice, does not threaten the integrity of the adjacent main blood vessel, and wherein main vessel patency is rapidly restored. A need also exists for such an

apparatus which may be used to seal off and occlude a peripheral vessel feeding an aneurysm or for other reasons, e.g. the preparation of a vein for use as an arterial bypass section without invasive surgery.

The present invention as characterized in claim 1 provides an elongated occluding balloon catheter for delivering an occluding agent. Preferred embodiments are further specified in the dependent claims

The reduced leakage of occluding agent back into the main vessel decreases the risk of vessel blockage and/or stenosis and resultant tissue ischemia. Patient comfort is increased and cost of the intensive care treatment is reduced by the shortened time and reduction of exposure to the occluding agent.

Figure 1 is a perspective view of one embodiment of the balloon catheter for occluding aneurysms or blood vessels of the invention;

Figure 2 is an end cross section view of the catheter body lumens of the catheter of Figure 1;

Figure 3 is a partial side cross section view of the catheter of Figure 1;

Figure 4A is an end cross section view of the catheter of Figure 3;

Figure 4B is an end cross section view of a variation of the balloon structure and delivery exit port of the catheter of Figure 3;

Figure 5 is a partial, side cross section view of a further embodiment of the balloon structure and delivery exit port of the catheter of Figure 1;

Figure 6 is an end cross section view of the catheter of Figure 5;

Figure 7 is a partial, side cross section view of a further embodiment of the balloon structure;

Figure 8A is an end cross section view of the catheter of Figure 7 configured to deliver a two component occluding agent through adjacent exit ports;

Figure 8B is an end cross section view of the catheter of Figure 7 configured differently than Figure 8A;

Figure 8C is a further end cross section view of the catheter of Figure 7 depicting a further method of forming the balloon structure;

Figure 9 is a perspective view of a further embodiment of the balloon catheter of the invention;

Figure 10A is an end cross section view of the catheter of Figure 9 depicting the balloon structure supporting a single lumen and delivery exit port;

Figure 10B is an end cross section view of the catheter of Figure 9 depicting the balloon structure supporting a double lumen and delivery exit ports;

Figure 11A is an end cross section view of the catheter of Figure 9 having a single delivery lumen formed in the outer tube;

Figure 11B is an end cross section view of the catheter of Figure 9 having two delivery lumens formed in the outer tube;

Figures 12A - 16B are schematic illustrations of the

steps of occluding an aneurysm in an artery with the catheter embodiments of the present invention. Figure 17 is an alternative schematic side view illustration of the step of introducing a balloon catheter in relation to the opening of a branch vessel into a main blood vessel to effect the occlusion thereof in the manner of Figures 12A - 16B; and Figure 18 depicts an alternate distal end segment of the balloon catheter of the various embodiments having a permanently installed torque wire attached therein for allowing rotation of the balloon.

In the following description, the several alternative preferred embodiments share common features of the invention which are illustrated generally in Figure 1 and more specifically in other figures. The methods of using the various embodiments to deliver an occluding agent and form an occluding cast in an aneurysm chamber or a branch vessel are illustrated schematically in Figures 12 - 17. The term "occluding agent" as defined above is intended to encompass the various liquid and solid materials and devices described herein which solidify or set up in situ or which encourage the formation of thrombus which occlude the aneurysm chamber or branch vessel.

The balloon catheter 10 of the present invention includes a length of multi-lumen flexible tubing forming a catheter body 12 having a proximal end segment 14 and a distal end segment 16. The catheter body 12 is preferably formed with a plurality of axially extending, co-linear passageways or lumens, e.g. the three lumens 18, 20 and 22 depicted in the cross section view of Figure 2 (and in further views) which are coupled to structure in the proximal and distal segments to function as a balloon inflation/deflation lumen and/or a guide wire lumen and as one or two delivery lumens. In other words, the three lumens 18, 20 and 22 can be configured to deliver a single form of occluding agent in one embodiment of the invention or to deliver two separate components of an occluding agent, e.g. a catalyst and an active ingredient, in a further embodiment of the invention as described in greater detail hereafter. Alternatively, one lumen may be used to vent or aspirate through the opening to assist in filling the aneurysm chamber or branch vessel with the single occluding agent. Four lumens may also be provided to deliver the two components or perform other functions.

In any such configuration, the tubing of the catheter body 12 may be extruded from flexible plastic materials, e.g. thermoplastics, polyvinyl chlorides, polyethylenes, polyurethanes, polyesters, polypropylenes or the like as is well known in the balloon catheter art. The catheter body may be extruded or formed with a variety of lumen cross sections, including circular or elliptic lumens (as shown in Figure 6) or in a co-axial configuration (as described with reference to Figures 9 - 11) or with the pie-shaped lumens depicted in Figure 2.

As shown in Figure 2A, the lumens 18, 20 and 22

are separated by webs 19 and 21 and confined in an outer tube 13. Lumen 18 is larger in cross section in order to accommodate a guide wire 70 shown in cross section. Lumens 20 and 22 are oriented together on one side to facilitate their alternate employment as delivery or venting lumens in the various embodiments described below.

Returning to Figure 1, the lumens 18, 20 and 22 are coupled through a manifold 32 at the catheter body proximal end segment 14 to a catheter proximal end connector assembly 26. One of the lumens, e.g. lumen 18, is coupled through manifold 32 to a single lumen tube 30 and proximally terminates in a fitting 28 into the aperture 24 of which a guide wire 70 may be inserted. In one embodiment, the guide wire lumen 18 is not employed for any other function, although it may also be employed as the balloon inflation/deflation lumen in other embodiments.

The second lumen 20 is coupled through manifold 32 to a tube 42 which is coupled in turn to a valve adaptor 40. The second lumen 20 may be coupled internally to the balloon 52 to function in one embodiment as an inflation/deflation lumen when it is fitted to a source of pressurized fluid (not shown) attached at adaptor fitting 38. In another embodiment, the lumen 20 may be in communication with a further delivery exit or venting port 60' adjacent to delivery exit port 60 and employed to deliver a component of a two component occluding agent or as an aspiration and/or venting lumen.

The third lumen 22 is coupled through manifold 32 to a tube 50 which is coupled in turn to a valve adaptor 48. The third lumen 22 may be coupled internally at the proximal junction 54 of the balloon 52 with the lead body 12 through an elastic tube extension formed either inside or outside the balloon in various embodiments and extending along the outer wall of the balloon 52 to a delivery exit port 60 positioned midway down the length of the balloon 52. The third lumen 22 functions in all of the embodiments as a conduit for the delivery of a contrast medium or occluding agent, as described hereafter, upon positioning of the delivery exit port with respect to the opening of an aneurysm and inflation of the balloon 52. The third lumen may also be employed to introduce further catheters or devices into proximity with the opening 60 as described below.

The balloon catheter 10 terminates at its distal end junction 55 with a soft tip 34 and a tip aperture 36 through which the guide wire 70 may extend during introduction of the catheter 10 and positioning of the balloon 52 alongside the aneurysm. The distal end segment 16 of the catheter body 12 is also provided with first and second radiopaque markers 15 and 17 which are located with respect to the delivery exit port 60 to assist in aligning it to the opening of an aneurysm or a branch vessel during introduction and orientation of the distal end segment 16. Any of the well known techniques may be employed for arterial and venous introduction of the catheter 10, with or without use of a

surrounding introduction catheter (not shown) or the guide wire 70.

Turning now to Figures 3 and 4A, they depict in a partial side cross section view and end view, one embodiment of the construction of the balloon 52 in relation to the catheter body 12, its lumens 18, 20, 22, and the tube extension 56 leading to the delivery exit port 60. In this cross section view, the lumens 20 and 22 are filled proximal to the distal aperture 36 to isolate the lumens and allow their use to deliver occluding agent and inflation fluid respectively. The inflation/deflation lumen 20 terminates and is filled more distally within the distal junction 55 of the balloon 52 with the outer surface of the lead body 12. The portion of the inflation/deflation lumen 20 within the balloon 52 has a number of spaced inflation/deflation holes 58 through the outer wall of the catheter body 12 to the interior space of the balloon 52. Inflation and deflation of balloon 52 are accomplished by applying and withdrawing pressurized fluid to and from the lumen 20 through the valve adaptor 40 in a manner well known in the art.

The delivery lumen 22 is coupled by through hole 23 to the lumen 57 of delivery tube extension 56. The delivery tube extension 56 is formed of a flexible, thin walled tube cemented alongside the balloon 52 externally to the balloon wall for the full length of the balloon 52 between the proximal and distal junctions 54 and 55. The through hole 23 extends through the side wall of delivery tube extension 56, which is cemented proximally to itself and the surface of the catheter body distal end segment 14, and makes a communication between the delivery tube extension lumen 57 and delivery lumen 22. The delivery tube extension lumen 57 is stopped up or filled, or adhered to itself, at its distal end lumen 62 distal to the delivery exit port 60 formed in its external facing wall.

The guide wire lumen 18 is open through the soft tip 34, which is preferably tapered in a manner well known in the balloon catheter art, and distal aperture 36. Several of the balloon inflation/deflation holes 58 through the inflation/deflation lumen 20 are depicted. The balloon 52 is preferably formed of a radiation cross-linked polyolefin, e.g. polyethylene, which does not readily adhere to the occluding agent contacting it during delivery and formation of the occluding cast and attached to the exterior surface of the catheter body 12 by adhesive or thermal bonding or welding at the proximal and distal junctions 54 and 55 in a manner well known in the art of fabricating miniature balloon catheters. The delivery extension tube may be formed of a thin walled TEFLON® polyethylene tube and adhered to the balloon 52 by adhesive or thermal bonding or welding. The balloon and delivery extension tube may also be coated with a release agent when fabricated of certain materials more prone to stick to the particular delivery agent.

Referring now to Figure 4B, it depicts, in an end cross section view conforming generally to Figure 4A, the above described features of the construction of the

distal end segment 16 and balloon 52 in accordance with a fabrication variation that may be employed in the first embodiment and is also depicted in Figure 7. In this variation, the delivery exit port 60 is formed in the outer wall of the balloon 52, and the delivery tube extension 56 is formed during balloon wall extrusion of the tubular shaped balloon. The exit port 60 is formed in the delivery tube extension 56 communicating with the lumen therein extending along the balloon 52 back to the proximal junction 54 and the through hole 23, as also shown in cross section in conjunction with the further embodiment of Figure 7.

The orientation of the delivery exit port 60 to the blood vessel opening is preferably determined through the use of the radiopaque markers 15 and 17 around the port 60 which may be observed during introduction and position through fluoroscopy. The proper seal afforded by the inflated balloon may be verified in the further steps of inflating the balloon 52, injecting a contrast medium through the delivery lumen 22 and exit port 60 after orienting the delivery exit port 60 to a trial position and observing the filling of the aneurysm as well as the absence of its leakage down the blood vessel lumen when it is properly oriented. Alternatively, a radiopaque tip probe may be introduced down the delivery lumen 22 and observed as it exits the exit port 60 and enters the aneurysm chamber or a pressure measurement may be taken.

In the embodiments described above, a liquid occluding agent is preferably introduced into the chamber of the aneurysm where it reacts with blood or tissue or solidifies to fill the space. Such occluding agents may include cross-linked collagen implant fibrils which may be mixed with contrast media and chemical buffers of the types described in the '718 patent. A liquid or paste collagen is available under the name HELIOSTAT®. A further liquid thrombin mixture is available under the name THROMBOSTAT®. Such thrombin and collagen including mixtures form an occluding cast by thrombus formation.

Other liquid, single component, occluding agents include methyl cyanoacrylate adhesives or 2-hydroxyethyl methylacrylate (HEMA) which set on contact with body fluids, e.g. of the type described in U.S. Patent No. 4,207,891 directed to occluding Fallopian tubes. In addition, liquid silicone rubber may be used.

Solid, single component, occluding agents may also be used in solid fibrous or particulate form that may be delivered through the vessel opening to form a solid mass of thrombus. The occluding agent is effective to coagulate blood around the fibers or particles and to form the thrombus mass within the aneurysm chamber or peripheral vessel to function as a solid occluding cast. Such occluding agents may also include one of the group of particulate compounds comprising polyvinyl alcohol (PVA), IVALON®, and GELL FOAM which are reactive to blood to coagulate it on contact, as described by Purdy in "Pre-Operative Embolization of

Cerebral Arteriovenous Malformations with Polyvinyl Alcohol Particles", AJNR 11:501-510, May/June, 1990.

It will also be appreciated that a solid occluding device or devices may be introduced through the lumen 22, out the delivery exit port 60 and through the adjoining vessel opening. For example, the wire coils described in the above referenced '437 patent may be so introduced, while the balloon 52 is inflated, to fill the aneurysm chamber or branch vessel from the vessel opening. Once introduced, a plurality of such coils entwine or catch on one another and the aneurysm or vessel side walls to provide acute fixation and encourage the formation of a mass of thrombus that forms the occluding cast.

A further feature of the invention is depicted in Figures 3 and 4A - 4B which allows the use of the balloon catheter 10 with certain occluding agents that are liquid until they are exposed to irradiating illumination of a frequency which causes the occluding agent to solidify. In this respect, a miniature probe or optical fiber 66 having a light diffuser or lens 68 at its distal end may be introduced down the lumen 22 (or lumen 20) and positioned to radiate light of the required frequency toward the opening 60. The catheter body 12 and balloon 52 may be transmissive of the frequency of radiation emitted, so that it falls upon occluding agent after it is delivered through the delivery exit port 60. Optionally, the balloon catheter 10 may be rotated within the vessel after the occluding agent is delivered to seal the aneurysm or branch vessel opening with the inflated balloon 52. The optical fiber 66 and diffuser 68 may be oriented to emit radiation through the catheter body distal end segment 16 and balloon 52 and toward the blood vessel opening to effect the curing of the occluding agent and the formation of the occluding cast. In Figures 4A and 4B, the optical fiber 66 and diffuser 68 are depicted preferably extending down the inflation/deflation lumen 20 and optionally down the delivery lumen 22, respectively. The optical fiber 66 and probe 68 may be extended down the inflation/deflation lumen 18 or the aspirating/venting lumens of the other embodiments as well.

In this variation, the occluding agent preferably comprises one of the group of light reactive compounds including urethane oligomer/(meth) acrylate monomer blends reactive to light in the ultraviolet range and particularly the compound Dymax 136-M[®] which is reactive to ultraviolet light of a frequency of 300-400 nanometers. Such compounds and light sources for their curing are described in the Dymax MD selector guide, Dymax data sheets and the Dymax 10M catalog.

As mentioned above, the lumens 18, 20 and 22 may be selectively configured to accommodate a two component occluding agent in a further embodiment of the invention depicted in partial cross section, elongated views of the distal end segment 16 in Figures 5 and 7, and in their respective end cross-section views in Figures 6 and 8. In order to use both of the lumens 20 and 22 as delivery lumens, it is preferable to provide a

further delivery exit port 60' and delivery tube lumen 61' extending from lumen 20 and through its wall in the catheter body distal end segment 16 closely adjacent to the delivery exit port 60 coupled to the delivery lumen 20.

Furthermore, it is preferable to employ the guide wire lumen 18 as the inflation/deflation lumen and to provide inflation/deflation holes 58 through its wall into the interior space of the balloon 52. And, in order to allow the guide wire to be extended through the elongated, tapered soft tip 34 and distal aperture 36, a self sealing valve 46 which is penetrable by the guide wire 70 is formed adjacent to the distal aperture 36. Such a self sealing valve 46 provides sufficient sealing against inflation pressure to allow inflation of the balloons 52 when the guide wire 70 is or is not extending through the valve 46 and may be of the type described in U.S. Patent No. 5,085,635 to Cragg.

In the embodiment of Figures 5 and 6, the lumen 20 is coupled to the lumen 61' of a further delivery tube extension 56' extending to the additional delivery exit port 60'. The further delivery tube extension 56' may be formed internal to the balloon or external to the balloon as a separate tube, as described above.

The balloon 52 depicted in Figures 6 and the preceding figures is generally cylindrical, encircling and extending along the catheter body distal end segment 16, so that the delivery exit port(s) are laterally displaced with the expanding balloon during inflation. In a further embodiment of the invention, the delivery exit port(s) 60, 60' are formed through the catheter body distal end segment 16, and an alternate balloon structure 52' is attached around only a major circumferential arc of the segment 16. The alternate balloon 52' configurations are depicted in a partial cross section, elongated view and end views of the distal end segment 16 in Figures 7, 8A, 8B and 8C.

In this embodiment and its depicted variations, alternate balloon 52' is formed along only a major circumferential section of the distal end segment 16 (Figures 8A and 8B) or a minor circumferential interior section of the tubular balloon is adhered along the outer surface of tube 13 (Figure 8C) so that the balloon 52' only inflates around the major circumference of the distal end segment 16. The delivery exit ports 60, 60' (or port 60) are formed through the outer tube 13 of the segment 16 in the minor circumferential section thereof in direct communication with the lumens 22 and 20 formed therein. The delivery exit ports 60 and 60', as shown in Figures 7 and 8C are thus directly made to the lumens 22 and 20, respectively, through the adhered balloon wall 52' and outer tube 13. The balloon 52' is inflated and deflated through the openings 58 made in the side wall of the catheter body distal end segment 16 to the lumen 18, which also functions as the guide wire lumen.

As also depicted in Figures 8A - 8C, the alternate balloon 52' can be formed to have alternate shapes

when inflated. Each of the balloons 52' are as roughly semi-circular and surrounding a major arc of the circumference of the distal end segment 16. The delivery exit ports 60 and 60' are formed in a minor circumferential arc or section of the distal end segment 16 of the catheter body 12. In Figure 8B, the balloon 52' forms a U-shaped perfusion channel 53, when inflated, along its length opposite to the minor section where the exit ports 60 and 60' are located. This alternate shape depicted in Figure 8B allows the balloon 52' to form the perfusion channel 53 with the blood vessel wall through which blood may continue to flow after the balloon is inflated in the vessel. In the other balloon shapes described above, it may be desirable to employ a separate perfusion catheter bypassing the balloon catheter 10 to provide perfusion while the occluding cast forms.

In the two component embodiment and these alternate configurations, the first and second components of an occluding agent may be delivered through the lumens 20 and 22 to exit the ports 60 and 60' after the balloon is inflated to position the ports against the opening of the aneurysm chamber. Such delivery is depicted in Figure 8A. The components mix together inside the aneurysm chamber or lumen of the branch vessel, and the resulting reaction solidifies the components to form the occluding cast therein. One example of a two component occluding agent would be catalyzable polyester monomers or epoxies.

Figure 8B also depicts the alternative use of the balloon catheter to deliver a single component occluding agent of the various types described above along with aspiration and/or ventilation of the contents of the aneurysm chamber or branch vessel lumen. In Figure 8B, the lumen 20 and delivery exit port 60' may be coupled at the proximal end 26 connectors to an aspirator to initially aspirate the contents of a branch blood vessel or aneurysm after the blood vessel opening is sealed. In addition or alternatively, the lumen 20 may be coupled to operate as a venting lumen to allow the contents to flow out as the occluding agent is delivered into the blood vessel opening through the delivery lumen 22 and exit port 60.

The above described features of the balloon catheter 10 of this embodiment with the alternate balloon 52' may be employed with only a single delivery lumen 22 and exit port 60, in the fashion of Figures 3 and 4 described above, to deliver a single component occluding agent. In this regard, Figure 7 may represent such a cross section of only a single delivery lumen 22 and exit port 60.

All of the above described embodiments and variations thereof may be implemented in the co-axial tube, preferably integral balloon configuration for the catheter body 12, including the catheter body proximal and distal end segments 14 and 16, as depicted in Figures 9 - 11. In the co-axial tube embodiment, the inner tube 11 is surrounded by an outer tube 13 to form the interior guide wire lumen 18 and the inflation lumen 20. The

outer tube 13 can be fabricated to form the inflation balloon 52 integrally with it in the fashion disclosed in the Simpson-Roberts U.S. Patent No. 4,323,071 in a manner well known in the balloon catheter art.

The single delivery lumen 22 is formed as shown in Figures 10A and 11A in the outer tube 13 extending the full length of the catheter body 12 and in the outer wall of the balloon 52 by standard multi-lumen extrusion techniques. The single delivery exit port 60 is formed as depicted in Figure 10A in the outer membrane of the outer tube 13 by standard skiving or porting techniques also well known in the balloon catheter fabrication art.

Figures 10B and 11B depict the addition of the second delivery or venting lumen 22' leading to the second delivery exit or venting port 60' for the applications described above. Thus, in this embodiment, the catheter body 12 is provided with four lumens. The configuration and construction of the embodiments of Figures 9 - 11 allows the catheter body 12 and balloon 52 to be integrally formed simply and with a low profile with a plurality of lumens formed in the outer tube 13 and balloon 52. Moreover, by adhering the balloon interior surface to the inner tube 11, in the manner of the embodiment of Figure 8C, the shapes of balloon 52' of Figures 7 and 8 may be employed with the features of this embodiment.

Turning now to the methods of use of the embodiments described above, reference is made to Figures 12 through 16 which are illustrations of an aneurysm 80 in an artery 90 and the steps of introducing and positioning a deflated balloon catheter, inflating the catheter, delivering the occluding agent, forming the cast and withdrawing the balloon catheter. Aneurysm 80 is formed through vessel opening 82 as a thin walled chamber 84 defined by wall 86. Figure 17 is a single illustration of the inflation and delivery of the occluding agent into the opening of a branch vessel corresponding to Figure 14B to illustrate that the same steps of Figures 12 - 16 would be employed in that method.

The introduction of the balloon catheter 10 through the arterial or venous system may be preceded by the introduction of the guide wire 70 in any of the various approaches employed in PTCA or balloon angioplasty. Once the guide wire is positioned, the balloon catheter 10, with the balloon 52 deflated, may be introduced over the guide wire 70 as depicted in Figure 12A. Alternatively, the guide wire 70 may be permanently attached to or positioned in the guide wire lumen 18 extending out the distal tip aperture 36 and/or valve 46, and the assembly may be advanced through the blood vessels until the deflated balloon 52, 52' is positioned and inflated as depicted in Figure 13A. The end views of Figures 12B and 13B depict these steps of positioning the guide wire 70 through the main vessel lumen 94 alongside the vessel or aneurysm opening 82 and inflating the balloon 52, 52'.

A guide catheter and/or introducer shrouding the balloon catheter 10 may also be employed in the introduction procedure, as is well known in the art. The

progress of introduction is typically observed under fluoroscopy and aided by the earlier identification of the aneurysm or branch vessel by radiopaque media which persists during the introduction and positioning steps.

Figure 12A is thus a schematic side view illustration of the aneurysm in an artery and the positioning of a deflated balloon catheter distal end segment 16 in relation to the aneurysm opening in accordance with the invention. It will be understood that a separate venting catheter 88 may also be positioned alongside the balloon catheter 10 so that the contents of the aneurysm chamber 84 may be aspirated and/or vented out as occluding agent is delivered through the delivery exit port 60, if the balloon catheter 10 is not configured to provide internal aspirating/venting as described above. Moreover, it will be understood that the balloon catheter of the invention in its various embodiments may be employed with a perfusion catheter 96 of any known type placed alongside the balloon 52, or in the U-shaped perfusion channel of the balloon 52', opposite to the opening 82. The perfusion catheter 96 allows blood flow past the temporary obstruction of the blood vessel 90 during the procedure without affecting the seal of the balloon to the vessel wall 92 afforded by the inflated balloon.

In this regard, once the balloon 52, 52' is positioned, it is oriented by twisting the proximal catheter body segment 14 so that the delivery exit port(s) 60, 60' is oriented facing the opening 82. The balloon positioning and orientation is observed under fluoroscopy to align the radiopaque markers 15, 17 to the opening 82. Testing of the seal may be accomplished by a test inflation while contrast medium is delivered as described above and as shown in Figures 14A and 14B. The test inflation may be avoided if the occluding agent is mixed with contrast media as described above. Alternatively, pressure readings may be taken through the delivery lumen to determine the adequacy of the balloon seal.

After the orientation and inflation is deemed acceptable, the delivery of the occluding agent or device may be accomplished. As the occluding agent is delivered, it either reacts with or displaces the fluid and blood clots in the chamber 84 and makes contact with the chamber wall 86. The delivery is depicted in Figures 14A and 14B.

The fully delivered occluding agent forming an occluding cast is illustrated in Figures 15A and 15B. As described above, the delivered occluding agent may be solidified through a number of alternative operations to form the occluding cast 100. During such occlusion, blood flow through the perfusion catheter or channel 53 afforded by the balloon shape may be continued.

After occlusion is completed and the occluding cast 100 is formed, the balloon 52 may be deflated and withdrawn along the guide wire 70. The guide wire 70 may then be moved.

Optionally, the balloon catheter 10 may be rotated after the delivery step of Figures 14A and 14B to re-ori-

ent the delivery exit opening(s) 60, 60' away from the opening 82 and seal the vessel opening 82 with the exterior side wall of the inflated balloon 52.

In this regard, it is necessary that the balloon catheter 10 be torsionally rigid enough down the length of the catheter body 12 to transmit rotational torque applied manually at its proximal end segment 14 to its distal end segment 16 in order to rotate the inflated balloon. Increased torque transmission ability may also be useful in introducing the catheter 10 and orienting the delivery exit port(s) 60, 60' to the vessel opening 82. It may therefore be desirable to increase the torque by providing a coupling of the distal segment of guide wire 70 with the lumen 18 or exit port 36 to allow torque to be transmitted by the guide wire. Alternatively, the guide wire may be fixedly attached permanently in the catheter lumen 18.

Figure 18 depicts an alternate distal end segment of the balloon catheter of the embodiment of Figure 7 having a permanently installed torque wire 71 attached distally therein by adhesive 72 to the soft tip 34 for increasing the torque transfer for allowing rotation of the balloon. The distally attached twist wire 71 is otherwise free in lumen 18 and may be coupled proximally to a knob to allow it to be twisted by the physician in the manner well known in the art and disclosed in U.S. Patent No. 4,582,181. It will be understood that this feature may be implemented in all of the embodiments described above by substitution with the guide wire 70 and associated structure.

The above described method is applicable as well to the occlusion of branch vessels 102 through a branch vessel opening 104 into the main vessel 90. Figure 17 depicts the intermediate step of that process, that is delivering the occluding agent into the branch vessel 102 through the opening 104, corresponding to Figure 14A. Each of the preceding and subsequent steps illustrated in Figures 12 - 16 may be followed in practicing the invention for the occlusion of branch vessels involving any of the preferred embodiments of the invention.

While a number of preferred embodiments of the invention and variations thereof have been described in detail, other modifications and methods of using and medical applications for the same will be apparent to those of skill in the art. Accordingly, it should be understood that various applications, modifications, and substitutions may be made of equivalents without departing from the scope of the invention as defined in the appended claims.

PARTS LIST

aneurysm occluding balloon catheter 10
inner tube 11
catheter body 12
outer tube 13
catheter body proximal end segment 14
first radiopaque marker 15

catheter body distal end segment 16
 second radiopaque marker 17
 guide wire lumen 18
 webs 19 and 21
 inflation/deflation lumen 20
 delivery lumen 22
 additional delivery or venting lumen 22'
 through hole 23
 aperture 24
 catheter proximal end connector assembly 26
 fitting 28
 single lumen tube 30
 manifold 32
 soft tip 34
 distal aperture 36
 adaptor fitting 38
 valve adaptor 40
 tube 42
 self sealing, penetrable, distal tip valve 46
 valve adaptor 48
 tube 50
 balloon 52
 alternate configuration balloon 52'
 U-shaped perfusion channel 53
 proximal junction 54
 distal junction 55
 elastic delivery tube extension 56
 additional elastic delivery tube extension 56'
 delivery tube extension lumen 57
 inflation/deflation holes 58
 delivery exit port 60
 additional delivery exit or venting port 60'
 delivery tube lumen 61
 additional delivery tube or venting tube lumen 61'
 filled distal most extension tube lumen 62
 filled distal most delivery lumen 63
 filled distal most inflation/deflation lumen 64
 optical fiber 66
 light radiation diffuser 68
 guide wire 70
 twist wire 71
 adhesive 72
 aneurysm 80
 aneurysm opening 82
 aneurysm chamber 84
 aneurysm wall 86
 venting catheter 88
 main vessel 90
 main vessel wall 92
 main vessel lumen 94
 bypass catheter 96
 occluding cast 100
 branch vessel 102
 branch vessel opening 104

Claims

1. An elongated occluding balloon catheter (10) for

delivering an occluding agent through a vessel Opening (104, 82) in the vessel wall (92) of a main blood vessel (90) leading to a peripheral vessel (102) or an aneurysm (80) for occluding the lumen of the peripheral vessel (102) or the aneurysm chamber (84) comprising:

an elongated catheter body (12) having a proximal end segment (14) and a distal end segment (16) adapted to be introduced into the main blood vessel lumen (94) to dispose the distal end segment extending along either side of the opening (104, 82);

an inflatable balloon (52) located at the catheter body distal end segment (16);

an elongated inflation/deflation lumen (20) extending through said catheter body proximal and distal end segments (14, 16) in fluid communication with the interior of said inflatable balloon (52) through which fluid may be delivered and withdrawn for inflating and deflating the inflatable balloon (52); and

an occluding agent delivery lumen (22) extending through the catheter body proximal and distal end segments (14, 16) and having a delivery exit port (60) in communication therewith located in said distal end segment (16); characterized in that:

said delivery exit port (60) is located intermediate proximal and distal junctions (54, 55) of said inflatable balloon (52) with said catheter body distal end segment (16) and alongside said inflatable balloon (52) such that upon orientation of said delivery exit port (60) in axial alignment with said opening (104, 82), the inflation of said inflatable balloon (52) causes it to bear against the main vessel wall (92) and urge said delivery exit port (60) against said opening (104, 82) to effect an occluding agent delivery passageway through said delivery exit port (60) and said opening (104, 82) that substantially inhibits any delivery of the occluding agent into the main vessel lumen (94).

2. The occluding balloon catheter (10) of Claim 1, further characterized in that:

said inflatable balloon (52) is formed of a balloon wall extending along said catheter body distal end segment (16) between said proximal and distal junctions (54, 55) and extending around the circumference of said catheter body distal end segment (16), whereby said inflatable balloon (52) expands outwardly from said catheter body distal end segment (16) to bear against the vessel wall (92) on inflation thereof; and

a distal portion of said delivery lumen (22) is

formed on and extends along said balloon wall a predetermined distance from said proximal junction (54) to said delivery exit port (60) which faces away from said balloon wall, whereby inflation of said inflatable balloon (52) laterally urges said delivery exit port (60) against the opening (104, 82) in the vessel wall (92).

3. The occluding balloon catheter (10) of Claim 1 or 2, further characterized in that:

said catheter body (12) further comprises an inner elongated tube (11) having an inner lumen (18) and an outer elongated tube (13) surrounding said inner elongated tube (11) and forming said inflation/deflation lumen (20) in said proximal catheter body end segment (16) and said inflatable balloon (52) in said catheter body distal end segment (16).

4. The occluding balloon catheter (10) of Claim 1 or 2, further characterized in that:

said catheter body (12) further comprises an inner elongated tube (11) having an inner lumen (18) and an outer elongated tube (13) surrounding said inner elongated tube (11) and forming said inflation/deflation lumen (20) in said proximal catheter body end segment (16) and the balloon wall of said inflatable balloon (52) in said catheter body distal end segment (16); and

said occluding agent delivery lumen (22) is formed within said outer tube (11) and extends through said catheter body proximal and distal end segments (14, 16), terminating in said delivery exit port (60).

5. The occluding balloon catheter (10) of Claim 3 or 4, further characterized by:

a distal tip (34) of said catheter body distal segment end (16);

a guide wire (70) for providing over the wire guidance for advancing the catheter body distal end segment (16) through the main blood vessel (90) to an occluding location with respect to said vessel opening (104, 82);

aperture means (24) at the proximal end of said inner lumen (18) for introducing said guide wire (70) into said inner lumen (18); and

an opening at said distal tip 34 through which said guide wire (70) is extendable.

6. The occluding balloon catheter (10) of Claim 1, further characterized in that:

said inflatable balloon (52) is formed of a balloon wall extending along said catheter body distal end segment (16) between said proximal and distal junctions (54, 55) and extending around at least a major section of the circumference thereof, whereby said inflatable balloon (52) expands outwardly from said catheter body distal end segment (16) to bear against the vessel wall (92) on inflation thereof and displaces a minor section of the circumference of said catheter body distal end segment (16) against the main vessel wall (92); and

a distal portion of said delivery lumen (22) extends distally through said elongated catheter body alongside said balloon wall a predetermined distance from said proximal junction (54) to said delivery exit port (60) which faces away from said balloon wall, whereby inflation of said inflatable balloon (52) laterally urges said delivery exit port (60) against the opening (104, 82) in the vessel wall (92).

7. The occluding balloon catheter (10) of Claim 6, further characterized in that:

said inflatable balloon (52) is formed of a tubular balloon wall adhered to said catheter body distal end segment (16) between said proximal and distal junctions (54, 55) in said minor section of the circumference thereof to form an adhered balloon section incapable of expanding on inflation of said inflatable balloon (52); and

said delivery exit port (60) is formed in said adhered balloon section, whereby inflation of said inflatable balloon (52) laterally urges said delivery exit port (60) against the vessel opening (104, 82).

8. The occluding balloon catheter (10) of any of the preceding Claims 2 - 7, wherein said inflatable balloon (52) is further characterized by:

means for forming a perfusion channel (53) diametrically opposite to said delivery exit port (60) and extending longitudinally alongside said balloon wall and the adjacent main vessel wall (92), whereby the perfusion channel (53) is sealed from the vessel opening (104, 82) upon inflation of the inflatable balloon (52) to allow the perfusion of blood in the perfusion channel.

9. The occluding balloon catheter (10) of any of the preceding Claims 1 - 8, further characterized by:

means for aspirating the contents of the aneurysm chamber (84) or the peripheral vessel (102) through the vessel opening (104, 82)

attendant to the delivery of the occluding agent through the delivery lumen (22) and the delivery exit port (60).

10. The occluding balloon catheter (10) of any of the preceding Claims 1 - 8, further characterized by:

means for venting the contents of the aneurysm chamber (84) or the peripheral vessel (102) through the vessel opening (104, 82) attendant to the delivery of the occluding agent through the delivery lumen (22) and the delivery exit port (60).

11. The occluding balloon catheter (10) of any of the preceding Claims 1 - 8, further characterized by:

a further elongated lumen (22') extending through said catheter body proximal and distal end segments (14, 16) and having a further exit port (60') in communication therewith located in said distal end segment (16) adjacent said delivery exit port (60) and alongside said inflatable balloon (52) such that upon orientation of said delivery exit port (60) and said further exit port (60') in axial alignment with said vessel opening (104, 82), the inflation of said inflatable balloon (52) causes it to bear against the main vessel wall (92) and urge said delivery exit port (60) and said further exit port (60') against said vessel opening (104, 82).

12. The occluding balloon catheter (10) of any of the preceding Claims 1 - 8, further characterized by means for venting the contents of the aneurysm chamber (84) or the peripheral vessel (102) through the vessel opening (104, 82) attendant to the delivery of the occluding agent through the delivery lumen (22) and the delivery exit port (60), said venting means further comprising:

a further elongated lumen (22') extending through said catheter body proximal and distal end segments (14, 16) and having a further exit port (60') in communication therewith located in said distal end segment (16) adjacent said delivery exit port (60) and alongside said inflatable balloon (52) such that upon orientation of said delivery exit port (60) and said further exit port (60') in axial alignment with said vessel opening (104, 82), the inflation of said inflatable balloon (52) causes it to bear against the main vessel wall (92) and urge said delivery exit port (60) and said further exit port (60') against said vessel opening (104, 82), whereby the contents of the aneurysm chamber (84) or the peripheral vessel (102) may be vented through the vessel opening (104, 82) attendant to the

delivery of the occluding agent through the delivery lumen (22) and the delivery exit port (60).

13. The occluding balloon catheter (10) of any of the preceding Claims 1 - 8, further characterized by means for aspirating the contents of the aneurysm chamber (84) or the peripheral vessel (102) through the vessel opening (104, 82) attendant to the delivery of the occluding agent through the delivery lumen (22) and the delivery exit port (60), said aspirating means further comprising:

a further elongated lumen (22') extending through said catheter body proximal and distal end segments (14, 16) and having a further exit port (60') in communication therewith located in said distal end segment (16) adjacent said delivery exit port (60) and alongside said inflatable balloon (52) such that upon orientation of said delivery exit port (60) and said further exit port (60') in axial alignment with said vessel opening (104, 82), the inflation of said inflatable balloon (52) causes it to bear against the main vessel wall (92) and urge said delivery exit port (60) and said further exit port (60') against said vessel opening (104, 82), whereby the contents of the aneurysm chamber (84) or the peripheral vessel (102) may be aspirated through the vessel opening (104, 82) attendant to the delivery of the occluding agent through the delivery lumen (22) and the delivery exit port (60).

14. The occluding balloon catheter (10) of any of the preceding Claims 1 - 13, wherein the occluding agent is reactive to irradiating light for solidifying as a solid occluding cast, and further characterized by:

means for irradiating the occluding agent delivered through the vessel opening (104, 82) with light directed through the vessel opening (104, 82) sufficiently to effect solidification of the delivered occluding agent.

15. The occluding balloon catheter (10) of any of the preceding Claims 1 - 13, wherein the occluding agent has a liquid form for delivery through the vessel opening (104, 82), the occluding agent being reactive on exposure to form the solid occluding cast.

16. The occluding balloon catheter (10) of any of the preceding Claims 1 - 13, wherein the occluding agent has a solid particulate form for delivery through the vessel opening (104, 82), the occluding agent effective to coagulate blood around the particles and to form a solid thrombus mass within the

aneurysm chamber (84) or peripheral vessel (102) that is functional as a solid occluding cast.

17. The occluding balloon catheter (10) of any of the preceding Claims 1 - 13, wherein the occluding agent delivered through the vessel opening (104, 82) is a first component of a two component occluding agent that reacts with a second component to form an occluding cast, and said occluding balloon catheter (10) is further characterized by:

a further elongated lumen (22') extending through said catheter body proximal and distal end segments (14, 16) and having a further exit port (60') in communication therewith located in said distal end segment (16) adjacent said delivery exit port (60) and alongside said inflatable balloon (52) through which said second component of said occluding agent is delivered such that upon orientation of said delivery exit port (60) and said further exit port (60') in axial alignment with said vessel opening (104, 82), the inflation of said inflatable balloon (52) causes it to bear against the main vessel wall (92) and urge said delivery exit port (60) and said further exit port (60') against said vessel opening (104, 82) to allow delivery of the first and second components of the two component occluding agent through said delivery exit port (60) and said further exit port (60'), respectively, and through said vessel opening (104, 82), whereby the first and second components of the occluding agent mix together in the peripheral vessel (102) or the aneurysm chamber (84) to form the occluding cast therein.

18. The occluding balloon catheter (10) of any of the preceding Claims 14 - 17, further characterized in that:

said elongated catheter body (12) is capable of being rotated manually from said proximal end segment (14) to first rotate the inflated balloon (52) and align said delivery exit port(s) (60, 60') with the vessel opening (104, 82) to effect delivery of the occluding agent(s) therethrough and to then further rotate the inflated balloon (52) following delivery of the occluding agent, whereby the vessel opening (104, 82) is sealed from the blood vessel lumen (94) as the occluding agent forms the occluding cast.

19. The occluding balloon catheter (10) of any of the preceding Claims 1 - 13, wherein the occluding agent is further characterized by a compressible shaped member having an expanded form when unconfined and compressible for delivery through a confined space and through the vessel opening

(104, 82), and specifically a coil or coils of noble metals.

20. The occluding balloon catheter (10) of any of the preceding Claims 1, 2 and 7 - 19, further characterized by:

a distal tip (34) of said catheter body distal segment end (16);

a guide wire (70) for providing over the wire guidance for advancing the catheter body distal end segment (16) through the main blood vessel (90) to an occluding location with respect to said vessel opening (104, 82);

aperture means (24) at said proximal end segment of said catheter body for introducing said guide wire (70) into said inflation/deflation lumen (20); and

penetrable valve means (46) formed in said distal tip (34) and extending across said inflation/deflation lumen (20) penetrable by said guide wire (70) and sealing against said guide wire (70) extending distally therethrough to prevent loss of inflation fluid during over the wire introduction and positioning of said occluding balloon catheter (10).

21. The occluding balloon catheter (10) of any of the preceding Claims 1, 2 and 7 - 19, further characterized by:

a distal tip (34) of said catheter body distal segment end (16); and

a twist wire (71) for providing guidance of said catheter body distal end segment (16) into position with respect to a vessel opening (104, 82) and for providing torque transfer from said catheter body catheter end segment (14) to said distal tip (34) to provide rotational movement of said delivery exit port (60) into alignment with the vessel opening (104, 82).

Patentansprüche

1. Langgestreckter Verschluß-Ballonkatheter (10) zum Zuführen bzw. Fördern eines Verschlußwirkstoffs bzw. Okkludiermittels durch eine Gefäßöffnung (104,82) in der Gefäßwand (92) eines zu einem Nebenblutgefäß (102) führenden Hauptblutgefäßes (90) oder eines Aneurysmas (80) zum Verschließen des Lumens des Nebenblutgefäßes (102) oder der Aneurysmakammer (84), umfassend:

einen langgestreckten Katheterkörper (12) mit einem proximalen Endsegment (14) und einem distalen Endsegment (16), das in das Lumen (94) des Hauptblutgefäßes eingeführt zu wer-

den vermag, um das distale Endsegment längs jeder Seite der Öffnung (104,82) verlaufend anzuordnen,

einen am distalen Endsegment (16) des Katheterkörpers gelegenen, aufblasbaren bzw. entfaltbaren Ballon (52),

ein sich durch die proximalen und distalen Endsegmente (14,16) des Katheterkörpers erstreckendes, längliches Inflations-/Deflationslumen (20), das in Fluidverbindung mit dem Inneren des aufblasbaren bzw. entfaltbaren Ballons (52) steht und durch das Fluid zum Aufblasen bzw. Ablassen des aufblasbaren Ballons (52) eingebracht bzw. abgeführt werden kann, sowie

ein sich durch die proximalen und distalen Endsegmente (14,16) des Katheterkörpers erstreckendes Lumen (22) zum Fördern bzw. Zuführen eines Verschluswirkstoffs bzw. Okkludiermittels, mit einer mit diesem in Verbindung stehenden, im distalen Endsegment (16) gelegenen Förder-Austrittsöffnung (60), dadurch gekennzeichnet, daß die Förder-Austrittsöffnung (60) in der Mitte zwischen proximalen und distalen Anschlüssen (54,55) des aufblasbaren Ballons (52) mit dem distalen Endsegment (16) des Katheterkörpers und längs des aufblasbaren Ballons (52) gelegen ist, so daß bei Orientierung der Förder-Austrittsöffnung (60) in axiale Ausrichtung auf die Öffnung (104,82) durch das Entfalten des aufblasbaren Ballons bewirkt wird, daß dieser sich an die Hauptgefäßwand (92) anlegt und die Förder-Austrittsöffnung (60) gegen die Öffnung (104,82) drängt, um durch die Förder-Austrittsöffnung (60) und die Öffnung (104,82) einen Durchgang für die Förderung bzw. Zuführung des Verschluswirkstoffs herzustellen, der im wesentlichen jegliches Einführen des Verschluswirkstoffs in das Lumen (94) des Hauptblutgefäßes verhindert.

2. Verschuß-Ballonkatheter (10) nach Anspruch 1, ferner dadurch gekennzeichnet, daß

der aufblasbare Ballon (52) aus einer sich längs des distalen Endsegments (16) des Katheterkörpers zwischen dem proximalen bzw. distalen Anschluß (54 bzw. 55) und um den Umfang des distalen Endsegments (16) des Katheterkörpers herum erstreckenden Ballonwand gebildet ist, wobei sich der aufblasbare Ballon (52) beim Aufblasen bzw. Entfalten von dem distalen Endsegment (16) des Katheterkörpers nach außen hin ausdehnt, um sich an die Gefäßwand (92) anzulegen, und ein distaler Endabschnitt des Förderlumens (22) an der Ballonwand gebildet ist und sich an

dieser entlang über eine vorbestimmte Entfernung von dem proximalen Anschluß (54) zu der Förder-Austrittsöffnung (60) hin erstreckt, welche von der Ballonwand weg gerichtet ist, wobei durch Entfalten des aufblasbaren Ballons (52) die Förder-Austrittsöffnung (60) gegen die Öffnung (104,82) in der Gefäßwand (92) gedrängt wird.

3. Verschuß-Ballonkatheter (10) nach Anspruch 1 oder 2, ferner dadurch gekennzeichnet, daß

der Katheterkörper (12) ferner einen inneren langgestreckten Schlauch (11) mit einem inneren Lumen (18) sowie einen äußeren langgestreckten Schlauch (13) aufweist, welcher den inneren langgestreckten Schlauch (11) umgibt und ein Inflations-/Deflationslumen (20) in dem proximalen Katheterkörper-Endsegment (16)(bzw. 14) und dem aufblasbaren Ballon (52) in dem distalen Katheterkörper-Endsegment (16) bildet.

4. Verschuß-Ballonkatheter (10) nach Anspruch 1 oder 2, ferner dadurch gekennzeichnet, daß

der Katheterkörper (12) ferner einen inneren langgestreckten Schlauch (11) mit einem inneren Lumen (18) und einen äußeren langgestreckten Schlauch (13) aufweist, welches den inneren langgestreckten Schlauch (11) umgibt und ein Inflations-/Deflationslumen (20) in dem proximalen Katheterkörper-Endsegment (16)(bzw. 14) und der Ballonwand des entfaltbaren Ballons (52) in dem distalen Katheterkörper-Endsegment (16) bildet, und das Förderlumen (22) für den Verschluswirkstoff innerhalb des äußeren Rohrs (11) gebildet ist und sich durch die proximalen und distalen Katheterkörper-Endsegmente (14,16) erstreckt und in der Förder-Austrittsöffnung (60) endet.

5. Verschuß-Ballonkatheter (10) nach Anspruch 3 oder 4, ferner gekennzeichnet durch

eine distale Spitze (34) des distalen Katheterkörper-Endsegments (16), einen Führungsdraht (70), um über die Drahtführung eine Vorwärtsbewegung des distalen Katheterkörper-Endsegments (16) durch das Hauptblutgefäß (90) zu einer Verschlusstelle bezüglich der Gefäßöffnung (104,82) zu bewirken, ein Öffnungsmittel (24) am proximalen Ende des inneren Lumens (18) zum Einführen des Führungsdrahts (70) in das innere Lumen (18), und eine Öffnung an der distalen Spitze (34), durch

die der Führungsdraht (70) einführbar ist.

6. Verschluß-Ballonkatheter (10) nach Anspruch 1, ferner dadurch gekennzeichnet, daß

der aufblasbare Ballon (52) aus einer sich längs des distalen Katheterkörper-Endsegments (16) zwischen den proximalen und distalen Anschlüssen (54,55) und um mindestens einen größeren Abschnitt seines Umfangs erstreckenden Ballonwand gebildet ist, wobei sich der aufblasbare Ballon (52) bei seiner Entfaltung von dem distalen Katheterkörper-Endsegment (16) nach außen ausdehnt, um sich gegen die Gefäßwand (92) anzulegen, und einen kleineren Abschnitt des Umfangs des distalen Katheterkörper-Endsegments (16) gegen die Hauptblutgefäßwand (92) verschiebt, und ein distaler Abschnitt des Förderlumens (22) sich distal durch den länglichen Katheterkörper längs der Ballonwand über eine vorbestimmte Strecke von dem proximalen Anschluß (54) bis zu der von der Ballonwand weg gerichteten Förder-Austrittsöffnung (60) erstreckt, wobei durch eine Entfaltung des aufblasbaren Ballons (52) die Förder-Austrittsöffnung (60) seitlich gegen die Öffnung (104,82) in der Gefäßwand (92) gedrängt wird.

7. Verschluß-Ballonkatheter (10) nach Anspruch 6, ferner dadurch gekennzeichnet, daß

der aufblasbare Ballon (52) aus einer schlauchförmigen Ballonwand gebildet ist, die an dem distalen Katheterkörper-Endsegment (16) zwischen den proximalen und distalen Anschlüssen (54,55) an dessen kleinerem Umfangsabschnitt festgeklebt ist, um einen verklebten Ballonabschnitt zu bilden, der sich bei der Entfaltung des aufblasbaren Ballons (52) nicht aufweiten kann, und die Förder-Austrittsöffnung (60) in dem verklebten Ballonabschnitt ausgebildet ist, wobei durch die Entfaltung des aufblasbaren Ballons (52) die Förder-Austrittsöffnung (60) seitlich gegen die Gefäßöffnung (104,82) gedrängt wird.

8. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 2 - 7, wobei der aufblasbare Ballon (52) ferner gekennzeichnet ist durch

Mittel zur Bildung eines zur Förder-Austrittsöffnung (60) diametral entgegengesetzten und sich in Längsrichtung entlang der Ballonwand und in Nähe der Hauptgefäßwand (92) erstreckenden Perfusionskanals (53), wobei der Per-

fusionskanal (53) beim Entfalten des aufblasbaren Ballons (52) gegenüber der Gefäßöffnung (104,82) abgedichtet wird, um die Perfusion von Blut im Perfusionskanal zu ermöglichen.

9. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 bis 8, ferner gekennzeichnet durch

eine Einrichtung zum Ansaugen des Inhalts der Aneurysmakammer (84) oder des Nebenblutgefäßes (102) durch die Gefäßöffnung (104,82) als Begleitmaßnahme zum Fördern bzw. Zuführen des Verschlußwirkstoffs durch das Förderlumen (22) und die Förder-Austrittsöffnung (60).

10. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 bis 8, ferner gekennzeichnet durch

eine Einrichtung zum Abführen des Inhalts der Aneurysmakammer (84) oder des Nebenblutgefäßes (102) durch die Gefäßöffnung (104,82) als Begleitmaßnahme zum Fördern bzw. Zuführen des Verschlußwirkstoffs durch das Förderlumen (22) und die Förder-Austrittsöffnung (60).

11. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 bis 8, ferner gekennzeichnet durch

ein weiteres langgestrecktes Lumen (22'), das sich durch die proximalen und distalen Katheterkörper-Endsegmente (14,16) erstreckt und eine weitere, mit ihm in Verbindung stehende und in dem distalen Endsegment (16) in Nähe der Förder-Austrittsöffnung (60) und längs des aufblasbaren Ballons (52) gelegene Austrittsöffnung (60') aufweist, so daß bei Orientierung der Förder-Austrittsöffnung (60) und der weiteren Austrittsöffnung (60') in axiale Ausrichtung auf die Gefäßöffnung (104,82) durch das Entfalten des aufblasbaren Ballons (52) bewirkt wird, daß dieser sich an die Hauptgefäßwand (92) anlegt und die Förder-Austrittsöffnung (60) und die weitere Austrittsöffnung (60') gegen die Gefäßöffnung (104,82) gedrängt werden.

12. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 bis 8, ferner gekennzeichnet durch

eine Einrichtung zum Abführen des Inhalts der Aneurysmakammer (84) oder des Nebenblut-

gefäßes (102) durch die Gefäßöffnung (104,82) als Begleitmaßnahme zum Fördern bzw. Zuführen des Verschlußwirkstoffs durch das Förderlumen (22) und die Förder-Austrittsöffnung (60), wobei die Abführeinrichtung ferner umfaßt:

ein weiteres langgestrecktes Lumen (22'), das sich durch die proximalen und distalen Katheterkörper-Endsegmente (14,16) erstreckt und eine weitere, mit ihm in Verbindung stehende und in dem distalen Endsegment (16) in Nähe der Förder-Austrittsöffnung (60) und längsseits des aufblasbaren Ballons (52) gelegene Austrittsöffnung (60') aufweist, so daß bei Orientierung der Förder-Austrittsöffnung (60) und der weiteren Austrittsöffnung (60') in axiale Ausrichtung auf die Gefäßöffnung (104,82), durch das Entfalten des aufblasbaren Ballons (52) bewirkt wird, daß dieser an der Hauptgefäßwand (92) anliegt und die Förder-Austrittsöffnung (60) und die weitere Austrittsöffnung (60') gegen die Gefäßöffnung (104,82) gedrängt werden, wobei der Inhalt der Aneurysmakammer (84) oder des Nebenblutgefäßes (102) durch die Gefäßöffnung (104,82) als Begleitmaßnahme zum Fördern bzw. Zuführen des Verschlußwirkstoffs durch das Förderlumen (22) und die Förder-Austrittsöffnung (60) abgeführt werden kann.

13. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 bis 8, ferner gekennzeichnet durch

eine Einrichtung zum Ansaugen des Inhalts der Aneurysmakammer (84) oder des Nebenblutgefäßes (102) durch die Gefäßöffnung (104,82) als Begleitmaßnahme zum Fördern bzw. Zuführen des Verschlußwirkstoffs durch das Förderlumen (22) und die Förder-Austrittsöffnung (60), wobei die Ansaugereinrichtung ferner umfaßt:

ein weiteres langgestrecktes Lumen (22'), das sich durch die proximalen und distalen Katheterkörper-Endsegmente (14,16) erstreckt und eine weitere, mit ihm in Verbindung stehende und in dem distalen Endsegment (16) in Nähe der Förder-Austrittsöffnung (60) und längsseits des entfaltenen Ballons (52) gelegene Austrittsöffnung (60') aufweist, so daß bei Orientierung der Förder-Austrittsöffnung (60) und der weiteren Austrittsöffnung (60') in axiale Ausrichtung auf die Gefäßöffnung (104,82) durch das Entfalten des aufblasbaren Ballons (52) bewirkt wird, daß dieser an der Hauptgefäßwand (92) anliegt und die Förder-Austrittsöffnung (60) und die weitere Austrittsöffnung (60') gegen die Gefäßöffnung (104,82) gedrängt

werden, wobei der Inhalt der Aneurysmakammer (84) oder des Nebenblutgefäßes (102) durch die Gefäßöffnung (104,82) als Begleitmaßnahme zum Fördern bzw. Zuführen des Verschlußwirkstoffs durch das Förderlumen (22) und die Förder-Austrittsöffnung (60) angesaugt werden kann.

14. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 bis 13, wobei der Verschlußwirkstoff auf Lichtbestrahlung anspricht, um sich als fester Verschlußpfropfen zu verfestigen, und ferner gekennzeichnet ist durch

eine Einrichtung zum Bestrahlen des durch die Gefäßöffnung (104,82) geförderten bzw. zugeführten Verschlußwirkstoffs mit Licht, das in ausreichendem Maße durch die Gefäßöffnung (104,82) gerichtet wird, um eine Verfestigung des zugeführten Verschlußwirkstoffs zu bewirken.

15. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 bis 13, wobei der Verschlußwirkstoff eine flüssige Form zum Fördern bzw. Zuführen durch die Gefäßöffnung (104,82) aufweist, wobei der Verschlußwirkstoff auf Lichteinwirkung anspricht, um den festen Verschlußpfropfen zu bilden.

16. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 bis 13, wobei der Verschlußwirkstoff für die Einbringung durch die Gefäßöffnung (104,82) eine feste Partikelform aufweist, wobei der Verschlußwirkstoff wirksam ist, um Blut um die Partikel herum zum Gerinnen zu bringen und eine feste Thrombenmasse innerhalb der Aneurysmakammer (84) oder des Nebenblutgefäßes (102) zu bilden, welche als fester Verschlußpfropfen fungiert.

17. Verschluß-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 bis 13, wobei der durch die Gefäßöffnung (104,82) zugeführte Verschlußwirkstoff eine erste Komponente eines Zweikomponenten-Verschlußwirkstoffs ist, der mit einer zweiten Komponente reagiert, um einen Verschlußpfropfen zu bilden, und der Verschluß-Ballonkatheter (10) ferner gekennzeichnet ist durch:

ein weiteres langgestrecktes Lumen (22'), das sich durch die proximalen und distalen Katheterkörper-Endsegmente (14,16) erstreckt und eine weitere, mit ihm in Verbindung stehende und in dem distalen Endsegment (16) in Nähe der Förder-Austrittsöffnung (60) und längsseits des aufblasbaren Ballons (52) gelegene Austrittsöffnung (60') aufweist, durch welche die

zweite Komponente des Verschlusswirkstoffs zugeführt wird, so daß bei Orientierung der Förder-Austrittsöffnung (60) und der weiteren Austrittsöffnung (60') in axiale Ausrichtung auf die Gefäßöffnung (104,82) durch das Entfalten des aufblasbaren Ballons (52) bewirkt wird, daß dieser sich an die Hauptgefäßwand (92) anlegt und die Förder-Austrittsöffnung (60) und die weitere Austrittsöffnung (60') gegen die Gefäßöffnung (104,82) gedrängt werden, um die Förderung bzw. Zuführung der ersten und zweiten Komponenten des Zweikomponenten-Verschlusswirkstoffs durch die Förder-Austrittsöffnung (60) bzw. die weitere Austrittsöffnung (60') sowie durch die Gefäßöffnung (104,82) zu ermöglichen, wobei sich die erste und die zweite Komponente des Verschlusswirkstoffs miteinander in dem Nebenblutgefäß (102) oder in der Aneurysmakammer (84) vermischen, um darin den Verschlusspfropfen zu bilden.

18. Verschluss-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 14 - 17, ferner dadurch gekennzeichnet, daß

der langgestreckte Katheterkörper (12) von dem proximalen Endsegment (14) aus manuell gedreht zu werden vermag, um zunächst den aufgeblasenen Ballon (52) zu drehen und die Förder-Austrittsöffnung(en) (60,60') auf die Gefäßöffnung (104,82) auszurichten, um eine Förderung bzw. Zuführung des (der) Verschlusswirkstoffs (Verschlusswirkstoffe) durch diese zu bewirken, und sodann den aufgeblasenen Ballon nach dem Einbringen des Verschlusswirkstoffs weiter zu drehen, wodurch die Gefäßöffnung (104,82) gegenüber dem Blutgefäßlumen (94) abgedichtet wird, während der Verschlusswirkstoff den Verschlusspfropfen bildet.

19. Verschluss-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1 - 13, wobei der Verschlusswirkstoff ferner gekennzeichnet ist durch ein komprimierbares Formelement, das eine ausge dehnte Form besitzt, wenn es nicht eingeschlossen ist, und zum Fördern durch einen eingeschlossenen Raum und durch die Gefäßöffnung (104,82) komprimierbar ist, und insbesondere eine Spule oder Spulen aus Edelmetallen.

20. Verschluss-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1, 2 und 7 - 19, ferner gekennzeichnet durch

eine distale Spitze (34) des distalen Katheterkörper-Endsegments (16),
einen Führungsdraht (70), um über die Draht-

führung eine Vorwärtsbewegung des distalen Katheterkörper-Endsegments (16) durch das Hauptblutgefäß (90) zu einer Verschlussstelle bezüglich der Gefäßöffnung (104,82) zu bewirken,

Öffnungsmittel (24) an dem proximalen Endsegment des Katheterkörpers zum Einführen des Führungsdrahts (70) in das Inflations-/Deflationslumen (20), sowie

in der distalen Spitze (34) ausgebildete und sich quer über das Inflations-/Deflationslumen (20) erstreckende, durchsetzbare Ventilmittel (46), die von dem Führungsdraht (70) durchsetzbar sind und gegen den Führungsdraht (70) abdichten, der sich distal durch sie hindurch erstreckt, um einen Verlust von Fluid zum Aufblasen bzw. Entfalten während des Einführens und Positionierens des Verschluss-Ballonkatheters (10) über den Draht zu verhindern.

21. Verschluss-Ballonkatheter (10) nach einem der vorangehenden Ansprüche 1, 2 und 7 - 19, ferner gekennzeichnet durch

eine distale Spitze (34) des distalen Katheterkörper-Endsegments (16), sowie
einen Torsionsdraht (71), um eine Führung des distalen Katheterkörper-Endsegments (16) in eine Position bezüglich der Gefäßöffnung (104,82) und eine Drehmomentübertragung von dem (proximalen) Katheterkörper-Endsegment (14) zur distalen Spitze (34) vorzusehen, um eine Drehbewegung der Förder-Austrittsöffnung (60) in Ausrichtung auf die Gefäßöffnung (104,82) zu bewirken.

Revendications

1. Cathéter (10) à ballonnet d'obturation allongé destiné à délivrer un agent de bouchage dans une ouverture de vaisseau (104, 82) dans la paroi (92) d'un vaisseau sanguin principal (90) conduisant à un vaisseau périphérique (102) ou un anévrisme (80) pour boucher le lumen du vaisseau périphérique (102) ou la chambre d'anévrisme (84) comprenant :

un corps de cathéter allongé (12) ayant un segment d'extrémité proximal (14) et un segment d'extrémité distal (16) adaptés pour être introduits dans le lumen (94) du vaisseau sanguin principal afin de disposer le segment d'extrémité distal s'étendant le long de l'un ou l'autre côté de l'ouverture (104, 82) ;
un ballonnet gonflable (52) situé au niveau du segment d'extrémité distal (16) de corps de cathéter ;

un lumen (20) de gonflage/dégonflage allongé s'étendant à travers lesdits segments d'extrémité proximal et distal (14, 16) de corps de cathéter en communication fluïdique avec l'intérieur dudit ballonnet gonflable (52) à travers lequel du fluide peut être envoyé et retiré pour gonfler et dégonfler le ballonnet gonflable (52) ; et

un lumen (22) de distribution d'agent de bouchage s'étendant dans les segments d'extrémité proximal et distal (14, 16) de corps de cathéter et ayant un orifice de sortie de distribution (60) en communication avec ceux-ci situé dans ledit segment d'extrémité distal (16) ; caractérisé en ce que :

ledit orifice de sortie de distribution (60) est situé entre les jonctions proximale et distale intermédiaires (54, 55) dudit ballonnet gonflable (52) avec ledit segment d'extrémité distal (16) de corps de cathéter et le long dudit ballonnet gonflable (52) de telle sorte que lors de l'orientation dudit orifice de sortie de distribution (60) en alignement axial avec ladite ouverture (104, 82), le gonflage dudit ballonnet gonflable (52) fait qu'il vient s'appuyer contre la paroi (92) du vaisseau principal et presser ledit orifice de sortie de distribution (60) contre ladite ouverture (104, 82) afin de réaliser un passage de distribution d'agent de bouchage dans ledit orifice de sortie de distribution (60) et ladite ouverture (104, 82) qui supprime sensiblement toute distribution de l'agent de bouchage dans le lumen (94) du vaisseau principal.

2. Cathéter (10) à ballonnet d'obturation selon la revendication 1, caractérisé en outre en ce que :

ledit ballonnet gonflable (52) est formé d'une paroi de ballonnet s'étendant le long dudit segment d'extrémité distal (16) de corps de cathéter entre lesdites jonctions proximale et distale (54, 55) et s'étendant sur la circonférence dudit segment d'extrémité distal (16) de corps de cathéter, ledit ballonnet gonflable (52) se dilatant vers l'extérieur depuis ledit segment d'extrémité distal (16) de corps de cathéter pour venir s'appuyer contre la paroi de vaisseau (92) lors de son gonflage ; et une partie distale dudit lumen de distribution (22) est formée sur et s'étend le long de ladite paroi de ballonnet à une distance prédéterminée depuis ladite jonction proximale (54) jusqu'àudit orifice de sortie de distribution (60) qui se trouve en face de ladite paroi de ballonnet, le gonflage dudit ballonnet gonflable (52) pressant latéralement ledit orifice de sortie de distribution (60) contre l'ouverture (104, 82)

dans la paroi de vaisseau (92).

3. Cathéter (10) à ballonnet d'obturation selon la revendication 1 ou 2, caractérisé en outre en ce que :

ledit corps de cathéter (12) comprend en outre un tube allongé intérieur (11) ayant un lumen intérieur (18) et un tube allongé extérieur (13) entourant ledit tube allongé intérieur (11) et formant ledit lumen de gonflage/dégonflage (20) dans ledit segment d'extrémité proximal (14) de corps de cathéter et dans ledit ballonnet gonflable (52) dans ledit segment d'extrémité distal (16) de corps de cathéter.

4. Cathéter (10) à ballonnet d'obturation selon la revendication 1 ou 2, caractérisé en ce que :

ledit corps de cathéter (12) comprend en outre un tube allongé intérieur (11) ayant un lumen intérieur (18) et un tube allongé extérieur (13) entourant ledit tube allongé intérieur (11) et formant ledit lumen de gonflage/dégonflage (20) dans ledit segment d'extrémité proximal (16) de corps de cathéter et la paroi de ballonnet dudit ballonnet gonflable (52) dans ledit segment d'extrémité distal (16) de corps de cathéter ; et

ledit lumen de distribution (22) d'agent de bouchage est formé à l'intérieur dudit tube extérieur (11) et s'étend dans lesdits segments d'extrémité proximal et distal (14, 16) de corps de cathéter, débouchant dans ledit orifice de sortie de distribution (60).

5. Cathéter (10) à ballonnet d'obturation selon la revendication 3 ou 4, caractérisé en outre par :

une pointe distale (34) dudit segment d'extrémité distal (16) de corps de cathéter ; un fil de guidage (70) pour fournir le guidage du fil afin d'avancer le segment d'extrémité distal (16) de corps de cathéter dans le vaisseau sanguin principal (90) jusqu'à un emplacement de bouchage par rapport à ladite ouverture de vaisseau (104, 82) ; des moyens d'ouverture (24) au niveau de l'extrémité proximale dudit lumen intérieur (18) pour introduire ledit fil de guidage (70) dans ledit lumen intérieur (18), et une ouverture au niveau de ladite pointe distale (34) à travers laquelle ledit fil de guidage (70) peut s'étendre.

6. Cathéter (10) à ballonnet d'obturation selon la revendication 1, caractérisé en outre en ce que :

- ledit ballonnet gonflable (52) est formé d'une paroi de ballonnet s'étendant le long dudit segment d'extrémité distal (16) de corps de cathéter entre lesdites jonctions proximale et distale (54, 55) et s'étendant autour d'au moins une section principale de sa circonférence, ledit ballonnet gonflable (52) se dilate vers l'extérieur depuis ledit segment d'extrémité distal (16) de corps de cathéter pour venir s'appuyer contre la paroi de vaisseau (92) lors du gonflage de celui-ci et déplace une petite section de la circonférence dudit segment d'extrémité distal (16) de corps de cathéter contre la paroi de vaisseau principal (92) ; et une partie distale dudit lumen de distribution (22) s'étend de manière distale dans ledit corps de cathéter allongé depuis ladite jonction proximale (54) jusqu'àudit orifice de sortie de distribution (60) qui fait face à ladite paroi de ballonnet, le gonflage dudit ballonnet gonflable (52) presse latéralement ledit orifice de sortie de distribution (60) contre l'ouverture (104, 82) dans la paroi de vaisseau (92).
7. Cathéter (10) à ballonnet d'obturation selon la revendication 6, caractérisé en outre en ce que :
- ledit ballonnet gonflable (52) est formé d'une paroi de ballonnet tubulaire collée audit segment d'extrémité distal (16) de corps de cathéter entre lesdites jonctions proximale et distale (54, 55) dans ladite petite section de sa circonférence pour former une section de ballonnet collée ne pouvant pas se dilater lors du gonflage dudit ballonnet gonflable (52) ; et ledit orifice de sortie de distribution (60) est formé dans ladite section de ballonnet collée, le gonflage dudit ballonnet gonflable (52) pressant latéralement ledit orifice de sortie de distribution (60) contre l'ouverture de vaisseau (104, 82).
8. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 2 à 7, dans lequel ledit ballonnet gonflable (52) est en outre caractérisé par :
- des moyens destinés à former un canal de perfusion (53) diamétralement opposé audit orifice de sortie de distribution (60) et s'étendant longitudinalement à côté de ladite paroi de ballonnet et de la paroi de vaisseau principal adjacente (92), le canal de perfusion (53) étant scellé depuis l'ouverture de vaisseau (104, 82) lors du gonflage du ballonnet gonflable (52) pour permettre la perfusion de sang dans le canal de perfusion.
9. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1 à 8, caractérisé en outre par :
- des moyens destinés à aspirer le contenu de la chambre d'anévrisme (84) ou du vaisseau périphérique (102) à travers l'ouverture de vaisseau (104, 82) associée à la distribution de l'agent de bouchage à travers le lumen de distribution (22) et l'orifice de sortie de distribution (60).
10. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1 à 8, caractérisé en outre par :
- des moyens destinés à purger le contenu de la chambre d'anévrisme (84) ou du vaisseau périphérique (102) à travers l'ouverture de vaisseau (104, 82) associée à la distribution de l'agent de bouchage à travers le lumen de distribution (22) et l'orifice de sortie de distribution (60).
11. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1 à 8, caractérisé en outre par :
- un autre lumen allongé (22') s'étendant à travers lesdits segments d'extrémité proximal et distal (14, 16) de corps de cathéter et ayant un autre orifice de sortie (60') en communication avec ceux-ci situé dans ledit segment d'extrémité distal (16) adjacent audit orifice de sortie de distribution (60) et le long dudit ballonnet gonflable (52) de telle sorte que lors de l'orientation dudit orifice de sortie de distribution (60) et dudit autre orifice de sortie (60') en alignement axial avec ladite ouverture de vaisseau (104, 82), le gonflage dudit ballonnet gonflable (52) fait qu'il vient s'appuyer contre la paroi de vaisseau principal (92) et presser ledit orifice de sortie (60) et ledit autre orifice de sortie (60') contre ladite ouverture de vaisseau (104, 82).
12. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1 à 8, caractérisé en outre par des moyens destinés à purger le contenu de la chambre d'anévrisme (84) ou du vaisseau périphérique (102) à travers l'ouverture de vaisseau (104, 82) associée à la distribution de l'agent de bouchage à travers le lumen de distribution (22) et de l'orifice de sortie de distribution (60), lesdits moyens de purge comprenant en outre :
- un autre lumen allongé (22') s'étendant à travers lesdits segments d'extrémité proximal et

distal (14, 16) de corps de cathéter et ayant un autre orifice de sortie (60') en communication avec ceux-ci situé dans ledit segment d'extrémité distal (16) adjacent audit orifice de sortie de distribution (60) et à côté dudit ballonnet gonflable (52) de telle sorte que lors de l'orientation dudit orifice de sortie de distribution (60) et dudit autre orifice de sortie (60') en alignement axial avec ladite ouverture de vaisseau (104, 82), le gonflage dudit ballonnet gonflable (52) fait qu'il vient s'appuyer contre la paroi de vaisseau principal (92) et presser ledit orifice de sortie de distribution (60) et ledit autre orifice de sortie (60') contre ladite ouverture de vaisseau (104, 82), le contenu de la chambre d'anévrisme (84) ou du vaisseau périphérique (102) pouvant être purgé à travers l'ouverture de vaisseau (104, 82) associée à la distribution de l'agent de bouchage à travers le lumen de distribution (22) et l'orifice de sortie de distribution (60).

13. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1 à 8, caractérisé en outre par des moyens pour aspirer le contenu de la chambre d'anévrisme (84) ou du vaisseau périphérique (102) à travers l'ouverture de vaisseau (104, 82) associée à la distribution de l'agent de bouchage par le lumen de distribution (22) et l'orifice de sortie de distribution (60), lesdits moyens d'aspiration comprenant en outre :

un autre lumen allongé (22') s'étendant à travers lesdits segments d'extrémité proximal et distal (14, 16) de corps de cathéter et ayant un autre orifice de sortie (60') en communication avec ceux-ci situé dans ledit segment d'extrémité distal (16) adjacent audit orifice de sortie de distribution (60) et à côté dudit ballonnet gonflable (52) de telle sorte que lors de l'orientation dudit orifice de sortie de distribution (60) et dudit autre orifice de sortie (60') en alignement axial avec ladite ouverture de vaisseau (104, 82), le gonflage dudit ballonnet gonflable (52) fait qu'il vient s'appuyer contre la paroi de vaisseau principal (92) et presser ledit orifice de sortie de distribution (60) et ledit autre orifice de sortie (60') contre ladite ouverture de vaisseau (104, 82), le contenu de la chambre d'anévrisme (84) ou du vaisseau périphérique (102) peut être aspiré à travers l'ouverture de vaisseau (104, 82) associée à la distribution de l'agent de bouchage à travers le lumen de distribution (22) et l'orifice de sortie de distribution (60).

14. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1 à 13,

dans lequel l'agent de bouchage est réactif à la lumière rayonnée pour se solidifier comme une coulée d'obturation solide, et caractérisé en outre par :

des moyens destinés à irradier l'agent de bouchage distribué à travers l'ouverture de vaisseau (104, 82) avec de la lumière dirigée dans l'ouverture de vaisseau (104, 82) suffisamment pour réaliser la solidification de l'agent de bouchage distribué.

15. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1 à 13, dans lequel l'agent de bouchage est sous forme liquide pour être distribué à travers l'ouverture de vaisseau (104, 82), l'agent de bouchage étant réactif à l'exposition pour former la coulée de bouchage solide.
16. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications 1 à 13, dans lequel l'agent de bouchage se trouve sous forme particulaire solide afin d'être distribué par l'ouverture de vaisseau (104, 82), l'agent de bouchage étant efficace pour faire coaguler le sang autour des particules et pour former un thrombus solide à l'intérieur de la chambre d'anévrisme (84) ou du vaisseau périphérique (102) qui fait office de coulée de bouchage solide.
17. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1 à 13, dans lequel l'agent de bouchage distribué par l'ouverture de vaisseau (104, 82) est un premier composant d'un agent de bouchage comprenant deux composants, qui réagit avec un second composant pour former une coulée de bouchage, et ledit cathéter à ballonnet d'obturation (10) est caractérisé en outre par :

un autre lumen allongé (22') s'étendant dans lesdits segments d'extrémité proximal et distal (14, 16) de corps de cathéter et ayant un autre orifice de sortie (60') en communication avec ceux-ci situé dans ledit segment d'extrémité distal (16) adjacent audit orifice de sortie de distribution (60) et à côté dudit ballonnet gonflable (52) à travers lequel ledit second composant dudit agent de bouchage est distribué de telle sorte que lors de l'orientation dudit orifice de sortie de distribution (60) et dudit autre orifice de sortie (60') en alignement axial avec ladite ouverture de vaisseau (104, 82), le gonflage dudit ballonnet gonflable (52) fait qu'il vient s'appuyer contre la paroi de vaisseau principal (92) et presse ledit orifice de sortie de distribution (60) et ledit autre orifice de sortie

(60') contre ladite ouverture de vaisseau (104, 82) pour permettre la distribution des premier et second composants de l'agent de bouchage comprenant deux composants à travers ledit orifice de sortie de distribution (60) et ledit 5
 autr orifice de sortie (60'), respectivement, et à travers ladite ouverture de vaisseau (104, 82) dans lequel les premier et second composants de l'agent de bouchage se mélangent dans le vaisseau périphérique (102) ou la chambre 10
 d'anévrisme (84) pour y former la coulée de bouchage.

18. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 14 à 17, caractérisé en outre en ce que :

ledit corps de cathéter allongé (12) peut être tourné manuellement depuis ledit segment d'extrémité proximal (14) pour d'abord faire 20
 tourner le ballonnet gonflé (52) et aligner ledit ou lesdits orifices de sortie de distribution (60, 60') avec l'ouverture de vaisseau (104, 82) pour effectuer la distribution du ou des agents de bouchage à travers celle-ci et pour ensuite 25
 faire tourner le ballonnet gonflé (52) après la distribution de l'agent de bouchage, l'ouverture de vaisseau (104, 82) étant scellée depuis le lumen de vaisseau sanguin (94) lorsque l'agent de bouchage forme la coulée de bouchage. 30

19. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1 à 13, dans lequel l'agent de bouchage est en outre caractérisé par un élément de forme compressible 35
 ayant une forme dilatée lorsqu'il n'est pas enfermé et compressible pour être distribué par un espace confiné et par l'ouverture de vaisseau (104, 82), et de manière spécifique une bobine ou des bobines en métaux nobles. 40

20. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1, 2 et 7 à 19, caractérisé en outre par :

une pointe distale (34) dudit segment d'extrémité distal (16) de corps de cathéter ;
 un fil de guidage (70) pour fournir le guidage du fil afin d'avancer le segment d'extrémité distal 50
 (16) de corps de cathéter par le vaisseau sanguin principal (90) jusqu'à un emplacement de bouchage par rapport à ladite ouverture de vaisseau (104, 82) ;
 des moyens d'ouverture (24) au niveau dudit segment d'extrémité proximal dudit corps de 55
 cathéter pour introduire ledit fil de guidage (70) dans ledit lumen de gonflage/dégonflage (20) ;
 et

des moyens de valve (46) pouvant pénétrer formés dans ladite pointe distale (34) et s'étendant dans ledit lumen de gonflage/dégonflage (20) dans lequel peut pénétrer ledit fil de guidage (70) et étant scellés contre ledit fil de guidage (70) s'étendant de manière distale à travers celui-ci pour empêcher toute perte de fluide de gonflage pendant l'introduction du fil et le positionnement dudit cathéter à ballonnet de bouchage (10).

21. Cathéter (10) à ballonnet d'obturation selon l'une quelconque des revendications précédentes 1, 2 et 7 à 19, caractérisé en outre par :

une pointe distale (34) dudit segment d'extrémité distal (16) de corps de cathéter ; et
 un fil torsadé (71) pour fournir un guide dudit segment d'extrémité distal (16) de corps de cathéter en position par rapport à une ouverture de vaisseau (104, 82) et pour fournir un transfert de couple depuis ledit segment d'extrémité proximal (14) de corps de cathéter vers ladite pointe distale (34) pour fournir un mouvement de rotation depuis ledit orifice de sortie de distribution (60) en alignement avec l'ouverture de vaisseau (104, 82).

Fig. 1

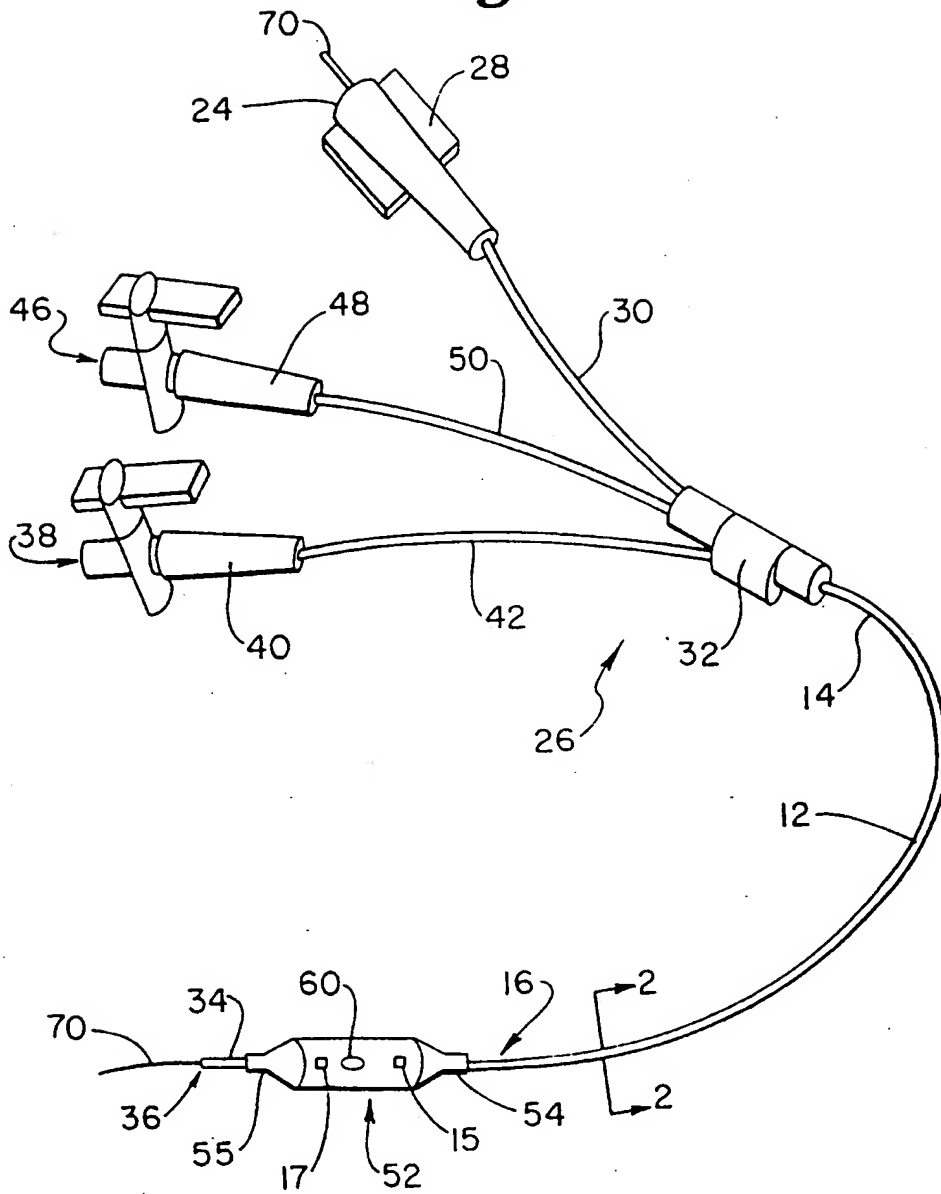


Fig. 2

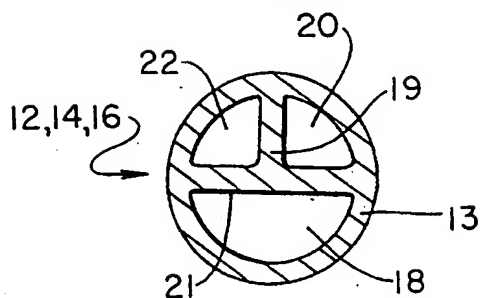


Fig. 3

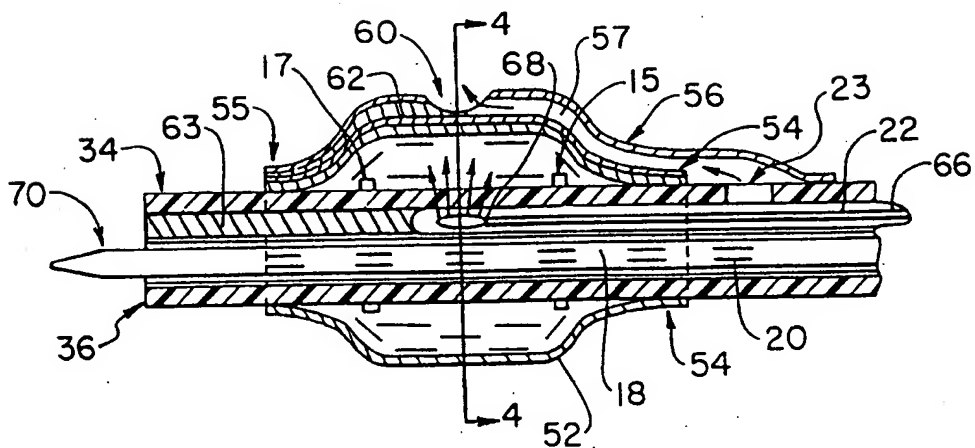


Fig. 4A

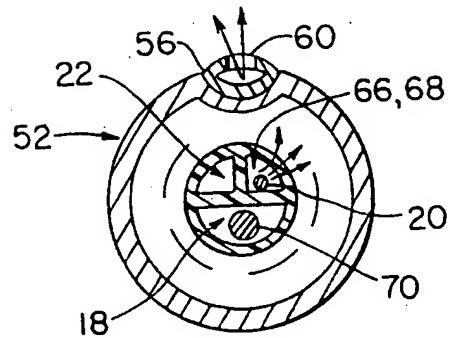


Fig. 4B

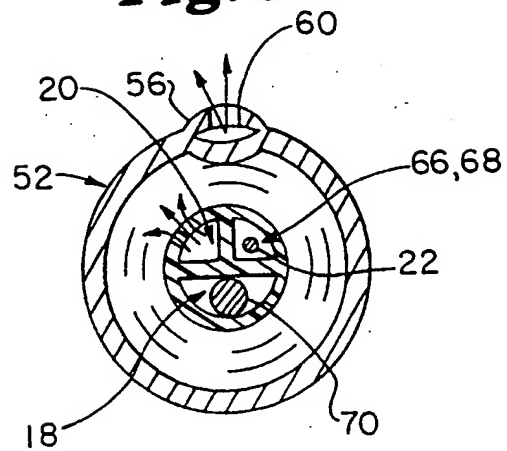


Fig.5

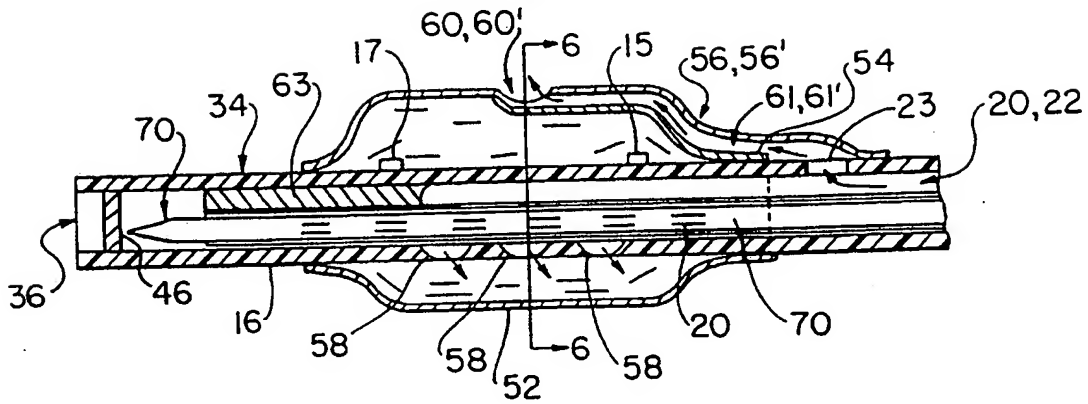


Fig.6

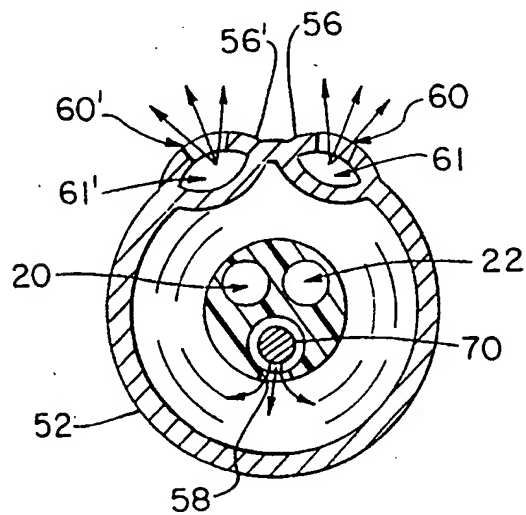


Fig. 7

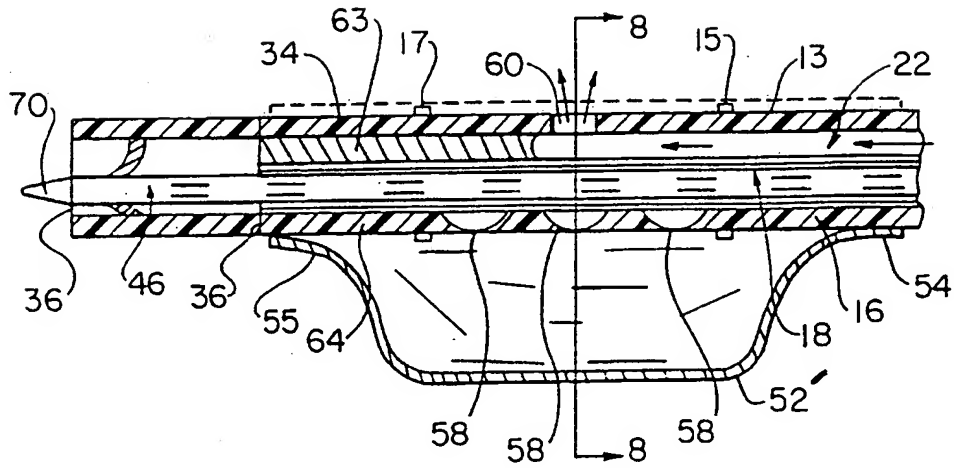


Fig. 8A

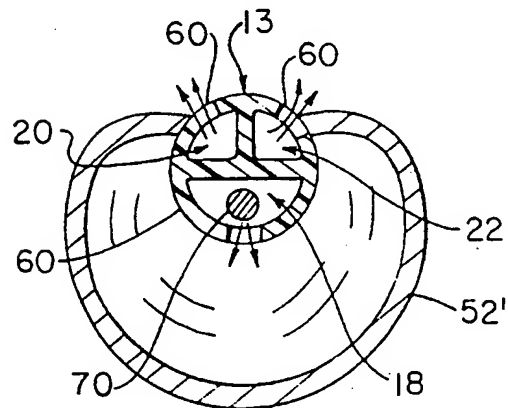


Fig. 8B

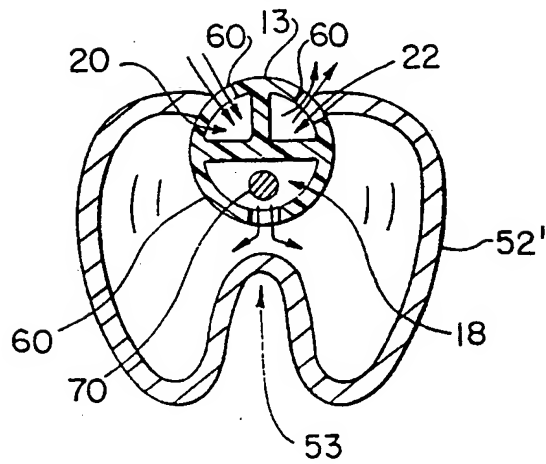


Fig.8C

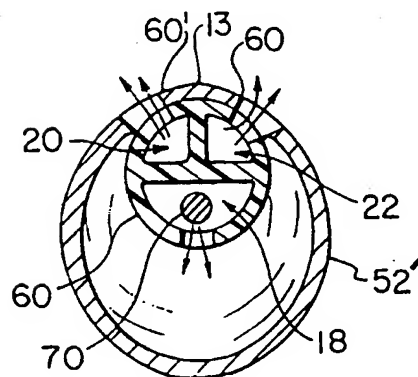


Fig. 9

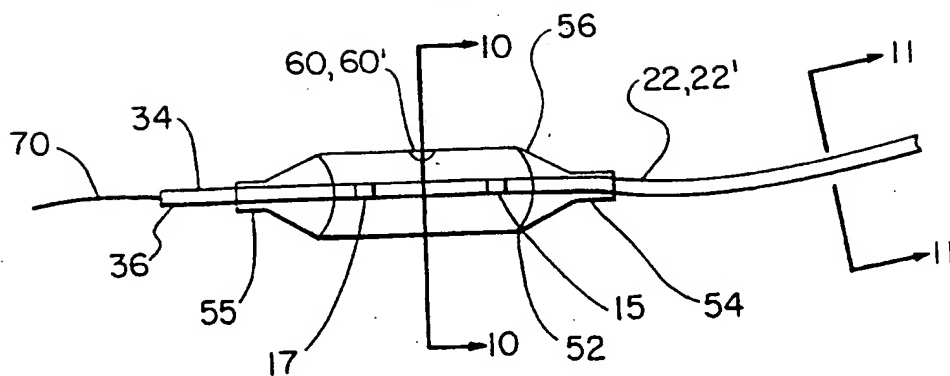


Fig. 10A

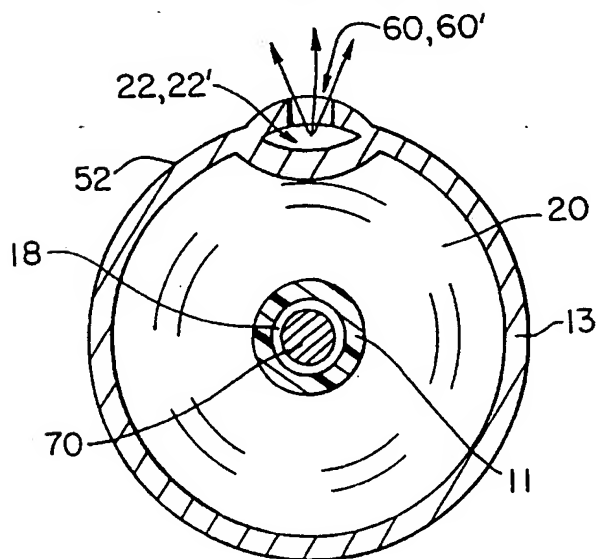


Fig.10B

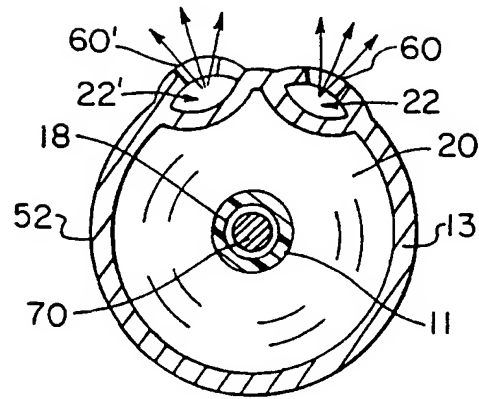


Fig.11A

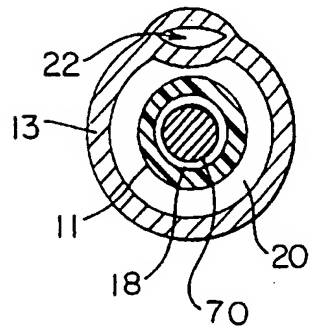


Fig.11B

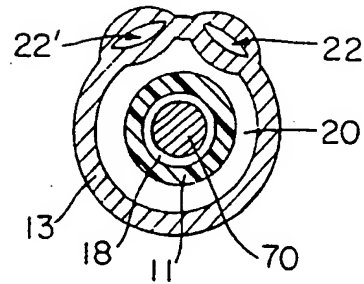


Fig.12A

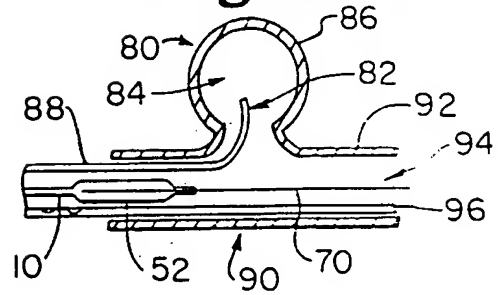


Fig.12B

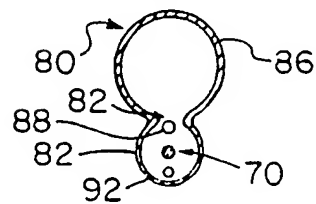


Fig.13A

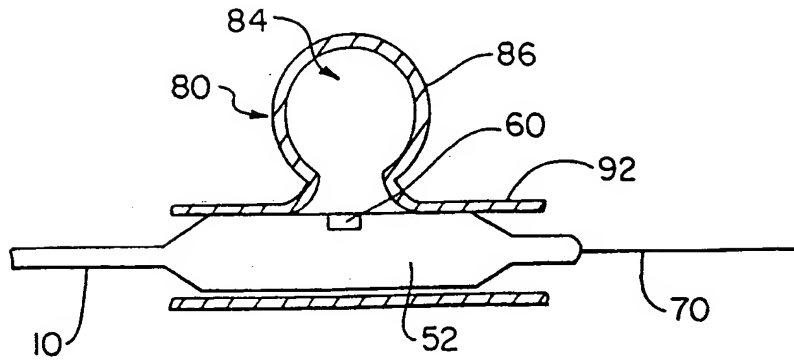


Fig.13B

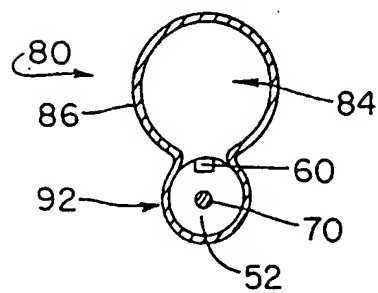


Fig.14A

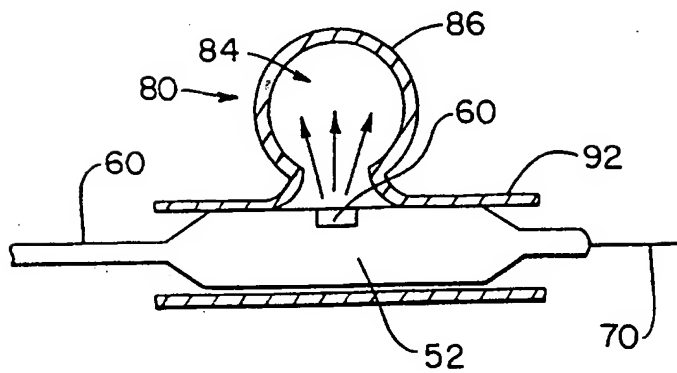


Fig.14B

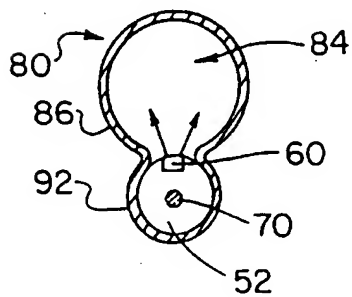


Fig.15A

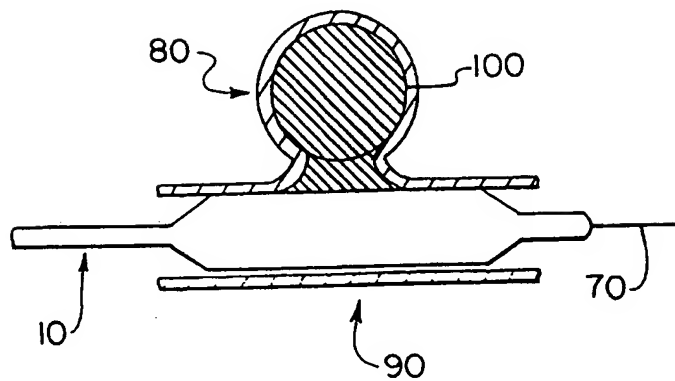


Fig.15B

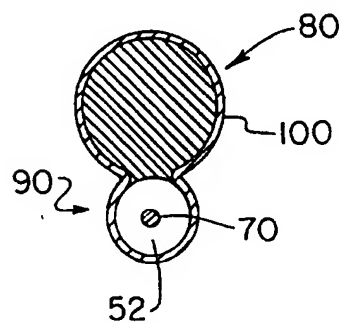


Fig.16A

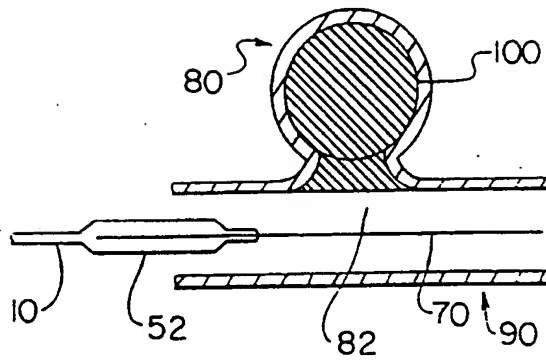


Fig.16B

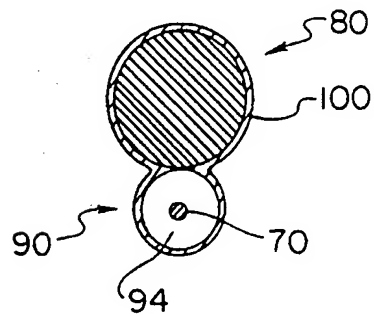


Fig. 17

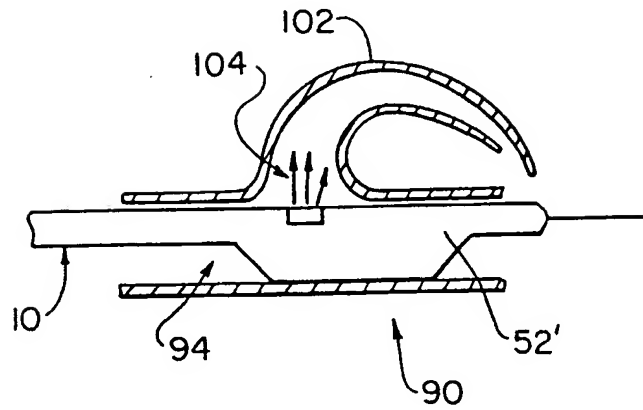


Fig. 18

